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PASSWORD:
TERMINAL (ENTER 1, 2, 3, OR ?):2

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=> FILE REGISTRY

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 20 JUL 2003 HIGHEST RN 551897-78-0 DICTIONARY FILE UPDATES: 20 JUL 2003 HIGHEST RN 551897-78-0

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

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Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=>
Uploading aryl nitrone for neuropathic pain.str

L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1 STR

G1 0, S

Structure attributes must be viewed using STN Express query preparation.

=> s l1

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100.0% PROCESSED 0 ITERATIONS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 0 TO 0
PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1

=> s l1 full

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100.0% PROCESSED 13 ITERATIONS 0 ANSWERS

0 ANSWERS

SEARCH TIME: 00.00.01

L3 0 SEA SSS FUL L1

=> fil marpat

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST 148.15 148.36

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FILE CONTENT: 1988-PRESENT (VOL 104 ISS 15-VOL 139 ISS03) (200300718ED)

MOST RECENT CITATIONS FOR PATENTS FROM FIVE MAJOR ISSUING AGENCIES (COVERAGE TO THESE DATES IS NOT COMPLETE):

US 6580000 17 JUN 2003 DE 20300703 26 JUN 2003 EP 1321471 25 JUN 2003 JP 2003173782 20 JUN 2003

WO 2003051918 26 JUN 2003

Structure search limits have been raised. See HELP SLIMIT for the new, higher limits.

=> s 11

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100.0% PROCESSED 223 ITERATIONS 1 ANSWERS SEARCH TIME: 00.00.03

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TI
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IN
     Jeon, Yoon T.; Gluchowski, Charles
PA
     Synaptic Pharmaceutical Corp., USA
SO
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     Preparation of 5-[4-(6-phenoxy-1-methyl-1H-benzimidazo-2-
    ylmethoxy)benzyl]thiazolidine-2,4-dione derivatives as antitumor agents
TN
     Shibata, Tomoyuki; Kurakata, Shinichi; Shimazaki, Naomi
PΑ
     Sankyo Company, Limited, Japan
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     PCT Int. Appl., 72 pp.
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LΑ
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AN
     138:255256 MARPAT
     Preparation of triazines as inhibitors of glycated protein-produced
TI
     induction of the signaling-assocd. inflammatory response in endothelial
IN
     Timmer, Richard T.; Alexander, Christopher W.; Pillarisetti, Sivaram;
     Saxena, Uday; Campbell, Karen A.
PA
     Reddy US Therapeutics, Inc., USA
     PCT Int. Appl., 396 pp.
SO
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AN
    138:238180 MARPAT
    Preparation of pyrazolopyridines for treatment of herpes infections
ΤI
IN
    Johns, Brian A.; Gudmundsson, Kristjan
PA
    Smithkline Beecham Corporation, USA
SO
    PCT Int. Appl., 124 pp.
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     138:187778 MARPAT
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     polymerase (PARP) inhibitors
IN
     Xu, Weizheng; Ferraris, Dana V.; Li, Jia-He; Kalish, Vincent J.
PA
     Guilford Pharmaceuticals Inc., USA
so
     PCT Int. Appl., 94 pp.
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     138:149044 MARPAT
ΤI
     Synergistic herbicidal compositions
IN
     Schaetzer, Juergen; Wenger, Jean; Hall, Roger Graham; Nebel, Kurt: Hole,
     Stephen
PA
     Syngenta Participations A.-G., Switz.
SO
     PCT Int. Appl., 47 pp.
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ΑN
     135:152622 MARPAT
ΤI
     Preparation of herbicidal substituted 1-phenyl-3-phenoxypropynes
IN
     Craig, Gerald Wayne; Eberle, Martin; Zeller, Martin; Bondy, Steven Scott;
     Comer, Daniel Dennis; Cheng, Soan; Penzotti, Julie Elizabeth; Grootenhuis,
     Peter Diederik Jan; Ehrler, Juerg
PA
     Syngenta Participations A.-G., Switz.
so
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AN
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    Color photographic silver halide material
IN
    Hagemann, Jorg; Haller, Jan; Helling, Gunter
PA
    Agfa-Gevaert N.V., Belg.
SO
    Eur. Pat. Appl., 39 pp.
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     Synergistic herbicides containing hydroxyphenylpyruvate dioxygenase
     inhibitors
IN
     Bieringer, Hermann; Van Almsick, Andreas; Hacker, Erwin; Willms, Lothar
PA
     Aventis Cropscience Gmbh, Germany
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     PCT Int. Appl., 41 pp.
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AN
     134:131317 MARPAT
     Preparation of 2-phenylaminobenzamides and analogs as MEK inhibitors for
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     the treatment of chronic pain
     Dixon, Alistair; Lee, Kevin; Pinnock, Robert Denham
IN
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     Warner-Lambert Company, USA
SO
     PCT Int. Appl., 132 pp.
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AN
     132:89504 MARPAT
TI
     Manufacture of amide-containing rodent repellent
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     Liu, Dongxue; Wei, Feng; Wang, Baorong; Wu, Yuchun; Fang, Hui
PA
     Shenyang Chemical Inst., Ministry of Chemical Industry, Peop. Rep. China
     Faming Zhuanli Shenqing Gongkai Shuomingshu, 36 pp.
SO
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AN
     130:168370 MARPAT
TI
     Preparation of hydantoin derivatives as farnesyl transferase inhibitors
IN
     Lee, Jin Ho; Koh, Jong Sung; Kim, Jong Hyun; Lee, Hyun Il; Jung, Won Hee;
     Ro, Seong Gu; Shin, You Seung; Kim, Sang Woong; Park, Ki Won; Kwak, Tae
     Hwan; Moon, Kyung Duk; Chung, Hyun Ho
PA
     LG Chemical Ltd., S. Korea
SO
     PCT Int. Appl., 129 pp.
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             THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
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AN
    129:230539 MARPAT
TI
    Preparation of herbicidal 2-cyano-3-phenyl-1,3-diones
IN
    Cain, Paul Alfred; Cramp, Susan Mary; Lambert, Claude; Wallis, Derek Ian;
    Yarwood, Thomas David; Little, Gillian Mary; Morris, John; Musil, Tibor;
    Pettit, Simon Neil; Smith, Philip Henry Gaunt; et al.
PA
    Rhone-Poulenc Agriculture Ltd., UK
SO
    U.S., 65 pp., Cont.-in-part of U.S. Ser. No. 304,482, abandoned.
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AN
     127:262678 MARPAT
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     Preparation of novel indoles and benzothiazoles for cloned human alpha 2
     receptors
TN
     Jeon, Yoon T.; Gluchowski, Charles
PΔ
     Synaptic Pharmaceutical Corp., USA
     PCT Int. Appl., 73 pp.
SO
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     ANSWER 14 OF 23 MARPAT COPYRIGHT 2003 ACS on STN
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AN
     125:107784 MARPAT
ΤI
     Pesticidal substituted diketonitriles
TN
     Cain, Paul A.; Chou, David T.; Da Silva, Themistocles D. J.; Gant, Daniel
     B.; Herman, Nancy D.
PA
     Rhone-Poulenc Inc., USA
SO
     Statutory Invent. Regist., 12 pp.
     CODEN: SRXXEV
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     125:33683 MARPAT
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     Aromatic amino ethers as pain relieving agents
IN
     Breault, Gloria Anne; Oldfield, John; Tucker, Howard; Warner, Peter
PA
     Zeneca Limited, UK
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     PCT Int. Appl., 140 pp.
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     ANSWER 16 OF 23 MARPAT COPYRIGHT 2003 ACS on STN
AN
     124:343326 MARPAT
     Preparation of substituted 4-aminopyrimido[5,4-d]pyrimidine antineoplastic
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IN
     Himmelsbach, Frank; Von rueden, Thomas
PA
     Dr. Karl Thomae Gmbh, Germany
so
     Ger. Offen., 39 pp.
     CODEN: GWXXBX
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     123:340138 MARPAT
     Preparation of heterocyclic-fused lactams which promote the release of
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     growth hormone
     Fisher, Michael H.; Mrozik, Helmut; Schoen, William R.; Shih, Thomas L.;
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     Wyvratt, Matthew J.
PA
     Merck and Co., Inc., USA
     PCT Int. Appl., 118 pp.
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     ANSWER 18 OF 23 MARPAT COPYRIGHT 2003 ACS on STN
AN
     123:313795 MARPAT
TI
     Preparation of ureidobenzaepinones as growth hormone release stimulants
IN
     Ok, Hyun O.; Schoen, William R.; Szumiloski, John
PA
     Merck and Co., Inc., USA
so
     PCT Int. Appl., 145 pp.
     CODEN: PIXXD2
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     ANSWER 19 OF 23 MARPAT COPYRIGHT 2003 ACS on STN
AN
     123:256544 MARPAT
ΤI
     Preparation of N-(oxobenzazepinyl)alkanamides as growth hormone release
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IN
     Schoen, William R.; Wyvratt, Matthew J., Jr.; Hodges, Paul J.
PA
     Merck and Co., Inc., USA
SO
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     123:228011 MARPAT
TI
     Preparation of N-(benzazepinonyl) alkanamides as growth hormone release
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IN
     Bochis, Richard J.; Hodges, Paul J.; Schoen, William R.; Wyvratt, Matthew
     J., Jr.
PΑ
     Merck and Co., Inc., USA
SO
     PCT Int. Appl., 185 pp.
     CODEN: PIXXD2
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     123:111666 MARPAT
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     Preparation of 1-aryl-2-cyano-3-cyclopropylpropane-1,3-diones as
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     Cain, Paul Alfred; Lambert, Claude; Cramp, Susan Mary; Little, Gillian
     Mary; Morris, John; Petit, Simon Neil; Smith, Philip Henry; Musil, Tibor
PA
     Rhone Poulenc Agriculture Ltd., UK
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    ANSWER 22 OF 23 MARPAT COPYRIGHT 2003 ACS on STN
L5
AN
     121:300926 MARPAT
ΤI
     (Heteroarylmethyl) indazole derivatives
IN
    Baker, Raymond; Kulagowski, Janusz Jozef; Leeson, Paul David; Smith,
    Adrian Leonard
PΑ
    Merck Sharp and Dohme Ltd., UK
    PCT Int. Appl., 48 pp.
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    ANSWER 23 OF 23 MARPAT COPYRIGHT 2003 ACS on STN
ΑN
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TT
    Induction of male sterility in crop plants with heterologous genes
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IN
    Hsu, Francis C.; Odell, Joan Tellefsen; Shen, Jennie Bih Jien
PA
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SO
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FILE COVERS 1907 - 21 Jul 2003 VOL 139 ISS 4 FILE LAST UPDATED: 20 Jul 2003 (20030720/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification. \cdot

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     Methods for the treatment of neuropathic pain by aryl nitrone compounds
     Waterbury, David; Wood, Paul L.; Khan, M. Amin; Upasani, Ravindra B.
TN
PA
     Centaur Pharmaceuticals, Inc., USA
SO
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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 20 JUL 2003 HIGHEST RN 551897-78-0 DICTIONARY FILE UPDATES: 20 JUL 2003 HIGHEST RN 551897-78-0

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

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L12 ANSWER 1 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-95-3 REGISTRY

FS 3D CONCORD

MF C23 H37 N O4

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

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1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 2 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN **452283-94-2** REGISTRY

CN 2-Propanamine, N-[[3,5-bis(1,1-dimethylethyl)-4-ethoxyphenyl]methylene]-2methyl-, N-oxide (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C21 H35 N O2

SR CA

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 3 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-93-1 REGISTRY

CN Benzenemethanamine, N-[[3,5-bis(1,1-dimethylethyl)-4-

(methoxymethoxy)phenyl]methylene]-, N-oxide (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C24 H33 N O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

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PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

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1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 4 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-92-0 REGISTRY

CN 2-Furanamine, N-[[3,5-bis(1,1-dimethylethyl)-4- (methoxymethoxy)phenyl]methylene]tetrahydro-, N-oxide (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C21 H33 N O4

SR CA

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1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 5 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-91-9 REGISTRY

CN Cyclohexanemethanamine, N-[[3,5-bis(1,1-dimethylethyl)-4-

(methoxymethoxy)phenyl]methylene]-, N-oxide (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C24 H39 N O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

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 CH_2-OMe
 CH_2-OMe

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 6 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-90-8 REGISTRY

CN Pentanoic acid, 2,6-bis(1,1-dimethylethyl)-4-formylphenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H30 O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 7 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN **452283-89-5** REGISTRY

CN Butanoic acid, 2,6-bis(1,1-dimethylethyl)-4-formylphenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H28 O3

SR CA

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 8 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN **452283-88-4** REGISTRY

CN Propanoic acid, 2-methyl-, 2,6-bis(1,1-dimethylethyl)-4-formylphenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H28 O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 9 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-87-3 REGISTRY

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[oxido(1,1,3,3-tetramethylbutyl)imino]methyl]-, acetate (ester) (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C25 H41 N O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

- 1 REFERENCES IN FILE CA (1947 TO DATE)
- 1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 10 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN **452283-86-2** REGISTRY

CN 2-Propanamine, N-[[3,5-bis(1,1-dimethylethyl)-4-methoxyphenyl]methylene]-2-methyl-, N-oxide (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H33 N O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 11 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN **452283-85-1** REGISTRY

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[oxido(phenylmethyl)imino]methyl]-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C22 H29 N O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

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1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 12 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-84-0 REGISTRY

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(1-methylethyl)oxidoimino]methyl]-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C18 H29 N O2

SR CA

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 13 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-83-9 REGISTRY

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylpropyl)oxidoimino]meth yl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H33 N O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

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1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 14 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN **452283-82-8** REGISTRY

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(2-hydroxy-1,1-dimethylethyl)oxidoimino]methyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H31 N O3

SR CA

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L12 ANSWER 15 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-81-7 REGISTRY

CN Pentanoic acid, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyl]phenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C24 H39 N O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

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1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 16 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-80-6 REGISTRY

FS 3D CONCORD

MF C20 H31 N O3

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LC STN Files: CA, CAPLUS, USPATFULL

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1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 17 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-79-3 REGISTRY

CN Butanoic acid, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyl]phenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C23 H37 N O3

SR CA

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 18 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 452283-78-2 REGISTRY

CN Propanoic acid, 2-methyl-, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyl]phenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C23 H37 N O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1947 TO DATE)

1 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 19 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273397-03-8 REGISTRY

CN Glycine, N-[[2,6-bis(1,1-dimethylethyl)-4-formylphenoxy]carbonyl]-, ethyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H29 N O5

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 20 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN . 273397-02-7 REGISTRY

FS 3D CONCORD

MF C21 H31 N O5

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 21 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273397-01-6 REGISTRY

CN Carbamic acid, butyl-, 2,6-bis(1,1-dimethylethyl)-4-formylphenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H31 N O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 22 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273397-00-5 REGISTRY

CN Carbamic acid, propyl-, 2,6-bis(1,1-dimethylethyl)-4-formylphenyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H29 N O3

SR CA

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 23 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273396-99-9 REGISTRY

CN Carbamic acid, ethyl-, 2,6-bis(1,1-dimethylethyl)-4-formylphenyl ester
(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C18 H27 N O3

SR · CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 24 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273396-96-6 REGISTRY

CN Methanethiol, [3-[[[3,5-bis(1,1-dimethylethyl)-4-(methoxymethoxy)phenyl]methylene]oxidoamino]propoxy]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C21 H35 N O4 S

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

$$P_{HS-CH_2-O-(CH_2)_3-N} = CH$$
 P_{U-t}
 $P_$

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 25 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273396-95-5 REGISTRY

FS 3D CONCORD

MF C22 H37 N O4 S

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 26 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273396-94-4 REGISTRY

CN 2-Propanamine, N-[[3,5-bis(1,1-dimethylethyl)-4-[(2-methoxyethoxy)methoxy]phenyl]methylene]-2-methyl-, N-oxide (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C23 H39 N O4

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

$$t-Bu-N$$
 CH
 $O-CH_2-O-CH_2-OMe$
 $t-Bu$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 27 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273396-93-3 REGISTRY

CN 2-Propanamine, N-[[3,5-bis(1,1-dimethylethyl)-4-(methoxymethoxy)phenyl]methylene]-2-methyl-, N-oxide (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C21 H35 N O3

SR CA

$$t-Bu-N$$
 CH
 $O-CH_2-OMe$
 $t-Bu$

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 28 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273396-92-2 REGISTRY

CN Glycine, N-[[2,6-bis(1,1-dimethylethyl)-4-[[(1,1dimethylethyl)oxidoimino]methyl]phenoxy]carbonyl]-, ethyl ester (9CI) (CA
INDEX NAME)

FS 3D CONCORD

MF C24 H38 N2 O5

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 29 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 273396-91-1 REGISTRY

CN .beta.-Alanine, N-[[2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyl]phenoxy]carbonyl]-, ethyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C25 H40 N2 O5

SR CA

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 30 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 273396-90-0 REGISTRY
- CN Carbamic acid, butyl-, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-
- dimethylethyl)oxidoimino]methyl]phenyl ester (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C24 H40 N2 O3
- SR CA
- LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 31 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 273396-89-7 REGISTRY
- CN Carbamic acid, propyl-, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-
- dimethylethyl)oxidoimino]methyl]phenyl ester (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C23 H38 N2 O3
- SR CA
- LC STN Files: CA, CAPLUS, USPATFULL

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 32 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN **273396-88-6** REGISTRY
- CN Carbamic acid, ethyl-, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyl]phenyl ester (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C22 H36 N2 O3
- SR CA

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

- L12 ANSWER 33 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 273396-87-5 REGISTRY
- FS 3D CONCORD
- MF C27 H37 N O6
- SR CA
- LC STN Files: CA, CAPLUS, USPATFULL

$$\begin{array}{c|c} \text{MeO} & \begin{array}{c|c} \text{O} & \\ \\ \end{array} \\ \text{OMe} & \begin{array}{c} \text{CH}_2 - \text{N} \\ \end{array} \\ \text{CH} \\ \end{array} \\ \begin{array}{c|c} \text{CH} \\ \end{array} \\ \begin{array}{c|c} \text{OAc} \\ \end{array}$$

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 34 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 273396-86-4 REGISTRY
- CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[(methyloxidoimino)methyl]-, acetate (ester) (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C18 H27 N O3
- SR CA
- LC STN Files: CA, CAPLUS, USPATFULL

$$Me = N = CH$$

$$OAc$$

$$t-Bu$$

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 35. OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 273396-85-3 REGISTRY
- CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyll]-, propanoate (ester) (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C22 H35 N O3
- SR CA
- LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 36 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 273396-84-2 REGISTRY
- FS 3D CONCORD
- MF C25 H30 F3 N O3
- SR CA
- LC STN Files: CA, CAPLUS, USPATFULL

$$CH_2-N$$
 CH_2-N
 C

- 2 REFERENCES IN FILE CA (1947 TO DATE)
- 2 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 37 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 273396-82-0 REGISTRY
- CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(2-hydroxy-1,1-
- dimethylethyl)oxidoimino]methyl]-, acetate (ester) (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C21 H33 N O4

$$\begin{array}{c|c} \text{Me} & \text{O} \\ & \parallel \\ \text{HO-CH}_2-\text{C-N} & \text{CH} \\ & \parallel \\ \text{Me} \\ & \text{OAc} \\ & \text{t-Bu} \\ \end{array}$$

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 38 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN **251086-09-6** REGISTRY

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyll-, acetate (ester) (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C21 H33 N O3

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPAT2, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1947 TO DATE)

3 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 39 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 243457-35-4 REGISTRY

CN Phenol, 4-[(hexyloxidoimino)methyl]-2,6-dimethyl- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C15 H23 N O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 40 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 198695-58-8 REGISTRY

CN Phenol, 4-[[(1,1-dimethylethyl)oxidoimino]methyl]-2,6-dimethoxy- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C13 H19 N O4

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1947 TO DATE)

3 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 41 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 165047-84-7 REGISTRY

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[oxido(1,1,3,3-tetramethylbutyl)imino]methyl]- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1,3,3-tetramethylbutyl)imino]methyl]-, N-oxide

FS 3D CONCORD

MF C23 H39 N O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

$$Me_3C-CH_2-C-N=CH$$

$$Me$$

$$Me$$

$$OH$$

$$OH$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1947 TO DATE)

2 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 42 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 151166-75-5 REGISTRY

CN Benzaldehyde, 3,5-bis(1,1-dimethylethyl)-4-(methoxymethoxy)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C17 H26 O3

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

$$t-Bu$$
 $O-CH_2-OMe$ $O+CH_2-OMe$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

13 REFERENCES IN FILE CA (1947 TO DATE)

13 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 43 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 124956-04-3 REGISTRY

CN Benzaldehyde, 3,5-bis(1,1-dimethylethyl)-4-[(2-methoxyethoxy)methoxy]-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H30 O4

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

$$t-Bu$$
 $O-CH_2-O-CH_2-CH_2-OMe$
 $Bu-t$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

6 REFERENCES IN FILE CA (1947 TO DATE)

6 REFERENCES IN FILE CAPLUS (1947 TO DATE)

L12 ANSWER 44 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

RN 93933-61-0 REGISTRY

CN Phenol, 4-[[(1,1-dimethylethyl)oxidoimino]methyl]-2,6-dimethyl- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Phenol, 4-[[(1,1-dimethylethyl)imino]methyl]-2,6-dimethyl-, N-oxide

FS 3D CONCORD

DR 165047-87-0

MF C13 H19 N O2

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 6 REFERENCES IN FILE CA (1947 TO DATE)
- 6 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 45 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 87133-21-9 REGISTRY
- CN Benzaldehyde, 4-(acetyloxy)-3,5-bis(1,1-dimethylethyl)- (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C17 H24 O3
- LC STN Files: BEILSTEIN*, CA, CAPLUS, TOXCENTER, USPAT7ULL (*File contains numerically searchable property data)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 4 REFERENCES IN FILE CA (1947 TO DATE)
- 4 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 46 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
- RN 29211-05-0 REGISTRY
- CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)oxidoimino]methyl]- (9CI) (CA INDEX NAME)
- OTHER CA INDEX NAMES:
- CN Nitrone, N-tert-butyl-.alpha.-(3,5-di-tert-butyl-4-hydroxyphenyl)- (8CI)
- CN Phenol, 2,6-bis(1,1-dimethylethyl)-4-[[(1,1-dimethylethyl)imino]methyl]-,
 N-oxide
- FS 3D CONCORD
- DR 165047-86-9
- MF C19 H31 N O2
- LC STN Files: BEILSTEIN*, CA, CAPLUS, CASREACT, IFICDB, IFIPAT, IFIUDB, TOXCENTER, USPATFULL
 - (*File contains numerically searchable property data)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 13 REFERENCES IN FILE CA (1947 TO DATE)
- 13 REFERENCES IN FILE CAPLUS (1947 TO DATE)
- L12 ANSWER 47 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN RN 25100-12-3 REGISTRY

```
CN
     Cyclohexanamine, N-hydroxy-, hydrochloride (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN
     Hydroxylamine, N-cyclohexyl-, hydrochloride (8CI)
OTHER NAMES:
CN
     Cyclohexylhydroxyamine hydrochloride
CN
     Cyclohexylhydroxylammonium chloride
CN
     N-Cyclohexylhydroxylamine hydrochloride
CN
     N-Hydroxycyclohexanamine hydrochloride
DR
     72762-44-8
     C6 H13 N O . Cl H
MF
                  BEILSTEIN*, CA, CAPLUS, CASREACT, CHEMCATS, CHEMLIST, CSCHEM,
LC
     STN Files:
       IFICDB, IFIPAT, IFIUDB, MSDS-OHS, SPECINFO, TOXCENTER, USPATFULL
         (*File contains numerically searchable property data)
     Other Sources:
                      EINECS**
         (**Enter CHEMLIST File for up-to-date regulatory information)
CRN
     (2211-64-5)
       ин-он
   HC1
              47 REFERENCES IN FILE CA (1947 TO DATE)
              47 REFERENCES IN FILE CAPLUS (1947 TO DATE)
L12 ANSWER 48 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
RN
     16649-50-6 REGISTRY
CN
     2-Propanamine, N-hydroxy-2-methyl- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Hydroxylamine, N-tert-butyl- (6CI, 8CI)
OTHER NAMES:
     2-Hydroxylamino-2-methylpropane
CN
     N-Hydroxy-tert-butylamine
CN
     N-t-Butylhydroxylamine
CN
     N-tert-Butylhydroxylamine
     tert-Butylhydroxylamine
CN
FS
     3D CONCORD
MF
     C4 H11 N O
CI
LC
     STN Files:
                  BEILSTEIN*, BIOSIS, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT,
       CHEMINFORMRX, IFICDB, IFIPAT, IFIUDB, MEDLINE, TOXCENTER, USPAT2,
       USPATFULL
         (*File contains numerically searchable property data)
HO-NH-Bu-t
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
             181 REFERENCES IN FILE CA (1947 TO DATE)
               2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
             181 REFERENCES IN FILE CAPLUS (1947 TO DATE)
               1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
L12 ANSWER 49 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
RN
     5100-34-5 REGISTRY
CN
     Propanoic acid, 3-isocyanato-, ethyl ester (9CI) (CA INDEX NAME)
```

```
OTHER CA INDEX NAMES:
    Propionic acid, 3-isocyanato-, ethyl ester (6CI, 7CI, 8CI)
OTHER NAMES:
     Ethyl 3-isocyanatopropanoate
CN
CN
     Ethyl 3-isocyanatopropionate
FS
     3D CONCORD
MF
     C6 H9 N O3
LC
     STN Files:
                  BEILSTEIN*, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, CSCHEM,
       IFICDB, IFIPAT, IFIUDB, TOXCENTER, USPAT2, USPATFULL
         (*File contains numerically searchable property data)
Eto-C-CH2-CH2-NCO
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
              54 REFERENCES IN FILE CA (1947 TO DATE)
              54 REFERENCES IN FILE CAPLUS (1947 TO DATE)
               4 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
L12 ANSWER 50 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
RN
     5080-22-8 REGISTRY
CN
     2-Propanamine, N-hydroxy- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Hydroxylamine, N-isopropyl- (6CI, 7CI, 8CI)
OTHER NAMES:
CN
     Isopropylhydroxylamine
CN
     N-Hydroxy-2-propanamine
CN
     N-Isopropylhydroxylamine
FS
     3D CONCORD
MF
     C3 H9 N O
CI
     COM
LC
     STN Files:
                 BEILSTEIN*, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, CHEMLIST,
       CIN, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, PIRA, PROMT, TOXCENTER,
       USPAT2, USPATFULL
         (*File contains numerically searchable property data)
                      EINECS**, NDSL**, TSCA**
         (**Enter CHEMLIST File for up-to-date regulatory information)
    NH-OH
H<sub>3</sub>C-CH-CH<sub>3</sub>
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
             140 REFERENCES IN FILE CA (1947 TO DATE)
               2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
             141 REFERENCES IN FILE CAPLUS (1947 TO DATE)
               5 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
L12 ANSWER 51 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
RN
     3970-21-6 REGISTRY
CN
     Ethane, 1-(chloromethoxy)-2-methoxy- (7CI, 8CI, 9CI)
                                                            (CA INDEX NAME)
OTHER NAMES:
     .beta.-Methoxyethoxymethyl chloride
CN
     1-(Chloromethoxy)-2-methoxyethane
CN
     2,5-Dioxahexyl chloride
     2-(Chloromethoxy)ethoxymethane
CN
```

```
2-Methoxyethoxymethyl chloride
CN
     Chloromethyl 2-methoxyethyl ether
CN
     Methoxyethoxymethyl chloride
FS
     3D CONCORD
MF
     C4 H9 Cl O2
LC
     STN Files:
                  BEILSTEIN*, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS,
       CHEMINFORMRX, CHEMLIST, CSCHEM, IFICDB, IFIPAT, IFIUDB, MEDLINE,
       SYNTHLINE, TOXCENTER, USPAT2, USPATFULL
          (*File contains numerically searchable property data)
     Other Sources:
                      EINECS**, NDSL**, TSCA**
         (**Enter CHEMLIST File for up-to-date regulatory information)
MeO-CH_2-CH_2-O-CH_2C1
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
             594 REFERENCES IN FILE CA (1947 TO DATE)
               2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
             595 REFERENCES IN FILE CAPLUS (1947 TO DATE)
               1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
L12
     ANSWER 52 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
RN
     2949-22-6 REGISTRY
     Acetic acid, isocyanato-, ethyl ester (6CI, 7CI, 8CI, 9CI)
CN
                                                                  (CA INDEX
     NAME)
OTHER NAMES:
CN
     Carbethoxymethyl isocyanate
CN
     Ethoxycarbonylmethyl isocyanate
CN
     Ethyl isocyanatoacetate
CN
     Isocyanatoacetic acid ethyl ester
FS
     3D CONCORD
MF
     C5 H7 N O3
CI
     COM
LC
     STN Files:
                  BEILSTEIN*, BIOBUSINESS, BIOSIS, CA, CAOLD, CAPLUS, CASREACT,
       CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, IFICDB, IFIPAT,
       IFIUDB, RTECS*, SPECINFO, SYNTHLINE, TOXCENTER, USPAT7, USPATFULL
         (*File contains numerically searchable property data)
                      EINECS**, NDSL**, TSCA**
         (**Enter CHEMLIST File for up-to-date regulatory information)
Eto-C-CH2-NCO
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
             286 REFERENCES IN FILE CA (1947 TO DATE)
               6 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
             286 REFERENCES IN FILE CAPLUS (1947 TO DATE)
               8 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
L12 ANSWER 53 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
```

(CA INDEX NAME)

CN

RN

CN

CN

CN

2211-64-5 REGISTRY

Cyclohexylhydroxylamine

N-Cyclohexylhydroxylamine

OTHER CA INDEX NAMES:

OTHER NAMES:

Cyclohexanamine, N-hydroxy- (9CI)

Hydroxylamine, N-cyclohexyl- (6CI; 7CI, 8CI)

CN N-Hydroxycyclohexanamine CN N-Hydroxycyclohexylamine FS 3D CONCORD MF C6 H13 N O CI COM LC STN Files: BEILSTEIN*, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CSCHEM, IFICDB, IFIPAT, IFIUDB, IPA, RTECS*, TOXCENTER, USPAT2, USPATFULL (*File contains numerically searchable property data) Other Sources: EINECS** (**Enter CHEMLIST File for up-to-date regulatory information) ин-он **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT** 188 REFERENCES IN FILE CA (1947 TO DATE) 189 REFERENCES IN FILE CAPLUS (1947 TO DATE) 40 REFERENCES IN FILE CAOLD (PRIOR TO 1967) L12ANSWER 54 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN RN 1620-98-0 REGISTRY CN Benzaldehyde, 3,5-bis(1,1-dimethylethyl)-4-hydroxy- (9CI) (CA INDEX NAME) OTHER CA INDEX NAMES: Benzaldehyde, 3,5-di-tert-butyl-4-hydroxy- (6CI, 7CI, 8CI) OTHER NAMES: 2,6-Di-tert-Butyl-4-formylphenol 3,5-Bis(1,1-dimethylethyl)-4-hydroxybenzaldehyde CN3,5-Di-tert-butyl-4-hydroxybenzaldehyde CN4-Formyl-2,6-di-tert-butylphenol CN 4-Hydroxy-3,5-di-tert-butylbenzaldehyde FS 3D CONCORD MFC15 H22 O2 CI COM LC STN Files: ANABSTR, BEILSTEIN*, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CSCHEM, HODOC*, IFICDB, IFIPAT, IFIUDB, RTECS*, SPECINFO, SYNTHLINE, TOXCENTER, USPAT2, USPATFULL (*File contains numerically searchable property data) Other Sources: EINECS** (**Enter CHEMLIST File for up-to-date regulatory information) Bu-t OHC OH t-Bu **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT** 585 REFERENCES IN FILE CA (1947 TO DATE) 6 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

L12 ANSWER 55 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

587 REFERENCES IN FILE CAPLUS (1947 TO DATE) 14 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

1122-60-7 REGISTRY RN CN Cyclohexane, nitro- (8CI, 9CI) (CA INDEX NAME) OTHER NAMES: CN Nitrocyclohexane FS 3D CONCORD MF C6 H11 N O2 CI COM LC ANABSTR, BEILSTEIN*, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, STN Files: CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CHEMSAFE, CSCHEM, DETHERM*, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2, HODOC*, HSDB*, IFICDB, IFIPAT, IFIUDB, MSDS-OHS, NIOSHTIC, RTECS*, SPECINFO, TOXCENTER, USPAT2, USPATFULL (*File contains numerically searchable property data) Other Sources: EINECS**, NDSL**, TSCA** (**Enter CHEMLIST File for up-to-date regulatory information) NO₂ **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT** 589 REFERENCES IN FILE CA (1947 TO DATE) 2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA 589 REFERENCES IN FILE CAPLUS (1947 TO DATE) 25 REFERENCES IN FILE CAOLD (PRIOR TO 1967) L12 ANSWER 56 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN 111-36-4 REGISTRY CNButane, 1-isocyanato- (9CI) (CA INDEX NAME) OTHER CA INDEX NAMES:

Isocyanic acid, butyl ester (6CI, 8CI) OTHER NAMES: CN1-Isocyanatobutane CNButyl isocyanate CNn-Butyl isocyanate FS 3D CONCORD MF C5 H9 N O CI COM LC AGRICOLA, ANABSTR, AQUIRE, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DETHERM*, DIPPR*, EMBASE, HSDB*, IFICDB, IFIPAT, IFIUDB, MEDLINE, MRCK*, MSDS-OHS, NAPRALERT, NIOSHTIC, PIRA, PROMT, RTECS*, SPECINFO, TOXCENTER, USPAT2, USPATFULL (*File contains numerically searchable property data) Other Sources: DSL**, EINECS**, TSCA** (**Enter CHEMLIST File for up-to-date regulatory information)

 $O = C = N - CH_2 - CH_2 - CH_2 - CH_3$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1680 REFERENCES IN FILE CA (1947 TO DATE)
103 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
1685 REFERENCES IN FILE CAPLUS (1947 TO DATE)
25 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L12 ANSWER 57 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN

```
RN
      110-78-1 REGISTRY
 CN
      Propane, 1-isocyanato- (9CI) (CA INDEX NAME)
 OTHER CA INDEX NAMES:
 CN
      Isocyanic acid, propyl ester (6CI, 7CI, 8CI)
 OTHER NAMES:
 CN
      1-Isocyanatopropane
 CN
      1-Propyl isocyanate
 CN
      n-Propyl isocyanate
 CN
      Propyl isocyanate
 FS
      3D CONCORD
 MF
      C4 H7 N O
 CI
      COM
 LC
      STN Files:
                   ANABSTR, BEILSTEIN*, BIOBUSINESS, CA, CAOLD, CAPLUS,
        CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CSCHEM, DETHERM*, HODOC*,
        IFICDB, IFIPAT, IFIUDB, NIOSHTIC, PIRA, RTECS*, SPECINFO, TOXCENTER,
        USPAT2, USPATFULL
          (*File contains numerically searchable property data)
                       DSL**, EINECS**, TSCA**
          (**Enter CHEMLIST File for up-to-date regulatory information)
 O = C = N - CH_2 - CH_2 - CH_3
 **PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
              834 REFERENCES IN FILE CA (1947 TO DATE)
               27 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
              840 REFERENCES IN FILE CAPLUS (1947 TO DATE)
                7 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
L12 ANSWER 58 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
 RN
      108-24-7 REGISTRY
      Acetic acid, anhydride (9CI)
                                     (CA INDEX NAME)
 OTHER CA INDEX NAMES:
      Acetic anhydride (8CI)
 OTHER NAMES:
      8: PN: US20030096775 PAGE: 17 claimed sequence
      8: PN: US20030105041 PAGE: 16 claimed sequence
      8: PN: US20030114401 PAGE: 17 claimed sequence
      8: PN: WO03040320 PAGE: 50 claimed sequence
      8: PN: WO03040321 PAGE: 52 claimed sequence
 CN
      8: PN: WO03040328 PAGE: 51 claimed sequence
      8: PN: WO03041657 PAGE: 51 claimed sequence
 CN
      8: PN: WO03042360 PAGE: 50 claimed sequence
 CN
      8: PN: WO03044163 PAGE: 67 claimed sequence
      8: PN: WO03046132 PAGE: 52 claimed sequence
      8: PN: WO03050246 PAGE: 51 claimed sequence
      98: PN: US20030113914 PAGE: 17 claimed sequence
      Acetic oxide
 CN
      Acetyl acetate
 CN
      Acetyl anhydride
 CN
      Acetyl ether
 CN
 CN
      Acetyl oxide
CN
      Ethanoic anhydride
 FS
      3D CONCORD
MF
      C4 H6 O3
CI
      COM
                   AGRICOLA, ANABSTR, AQUIRE, BEILSTEIN*, BIOBUSINESS, BIOSIS,
LC
      STN Files:
        BIOTECHNO, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS,
        CHEMINFORMRX, CHEMLIST, CHEMSAFE, CIN, CSCHEM, CSNB, DETHERM*, DIPPR*,
        EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2, GMELIN*, HODOC*,
        HSDB*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK*, MSDS-OHS, NIOSHTIC,
        PDLCOM*, PIRA, PROMT, RTECS*, SPECINFO, SYNTHLINE, TOXCENTER, TULSA,
```

ULIDAT, USPAT2, USPATFULL, VTB
 (*File contains numerically searchable property data)
Other Sources: DSL**, EINECS**, TSCA**
 (**Enter CHEMLIST File for up-to-date regulatory information)

Ac-O-Ac

```
**PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**
           13889 REFERENCES IN FILE CA (1947 TO DATE)
             350 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
           13935 REFERENCES IN FILE CAPLUS (1947 TO DATE)
              4 REFERENCES IN FILE CAOLD (PRIOR TO 1967)
L12 ANSWER 59 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN
RN
     107-30-2 REGISTRY
CN
     Methane, chloromethoxy- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Ether, chloromethyl methyl (6CI, 8CI)
OTHER NAMES:
CN
     .alpha.,.alpha.-Dichlorodimethyl ether
CN
     Chlorodimethyl ether
CN
     Chloromethoxymethane
CN
     Chloromethyl methyl ether
CN
     Methoxychloromethane
CN
     Methoxymethyl chloride
CN
     Methyl chloromethyl ether
CN
     Monochlorodimethyl ether.
CN
     Monochloromethyl methyl ether
FS
     3D CONCORD
MF
     C2 H5 C1 O
CI
     COM
                  BEILSTEIN*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT,
LC
       CAOLD, CAPLUS, CASREACT, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST,
       CHEMSAFE, CIN, CSCHEM, CSNB, DETHERM*, DIPPR*, EMBASE, ENCOMPLIT
       ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2, GMELIN*, HODOC*, HSDB*, IFICDB,
       IFIPAT, IFIUDB, IPA, MEDLINE, MRCK*, MSDS-OHS, NIOSHTIC, PDLCOM*, PROMT,
       RTECS*, SPECINFO, SYNTHLINE, TOXCENTER, ULIDAT, USPAT2, USPATFULL
         (*File contains numerically searchable property data)
     Other Sources:
                     EINECS**, NDSL**, TSCA**
         (**Enter CHEMLIST File for up-to-date regulatory information)
C1-CH2-O-CH3
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PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2679 REFERENCES IN FILE CA (1947 TO DATE)
132 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
2682 REFERENCES IN FILE CAPLUS (1947 TO DATE)
63 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L12 ANSWER 60 OF 60 REGISTRY COPYRIGHT 2003 ACS on STN RN 97-72-3 REGISTRY
CN Propanoic acid, 2-methyl-, anhydride (9CI) (CA INDEX NAME) OTHER CA INDEX NAMES:
CN Isobutyric anhydride (6CI, 7CI, 8CI) OTHER NAMES:
CN 2-Methylpropanoic anhydride
CN 2-Methylpropionic anhydride

CN Isobutanoic anhydride

CN Isobutyric acid anhydride

CN Isobutyryl anhydride

FS 3D CONCORD

MF C8 H14 O3

CI COM

LC STN Files: BEILSTEIN*, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, DETHERM*, HODOC*, HSDB*, IFICDB, IFIPAT, IFIUDB, MSDS-OHS, NIOSHTIC, SPECINFO, SYNTHLINE, TOXCENTER, USPAT2, USPATFULL

(*File contains numerically searchable property data)

Other Sources: DSL**, EINECS**, TSCA**

(**Enter CHEMLIST File for up-to-date regulatory information)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

744 REFERENCES IN FILE CA (1947 TO DATE)

7 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

744 REFERENCES IN FILE CAPLUS (1947 TO DATE)

12 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> sel rn 3-5, 24-27 E61 THROUGH E67 ASSIGNED

=> fil medl hcapl biosis uspatf
COST IN U.S. DOLLARS

SINCE FILE

ENTRY 104.16

TOTAL SESSION 392.18

FULL ESTIMATED COST

FILE 'MEDLINE' ENTERED AT 12:11:23 ON 21 JUL 2003

FILE 'HCAPLUS' ENTERED AT 12:11:23 ON 21 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'BIOSIS' ENTERED AT 12:11:23 ON 21 JUL 2003 COPYRIGHT (C) 2003 BIOLOGICAL ABSTRACTS INC.(R)

FILE 'USPATFULL' ENTERED AT 12:11:23 ON 21 JUL 2003 CA INDEXING COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

=> s e61-67

L13

L14

2 (273396-93-3/BI OR 273396-94-4/BI OR 273396-95-5/BI OR 273396-96 -6/BI OR 452283-91-9/BI OR 452283-92-0/BI OR 452283-93-1/BI)

=> dup rem 113
PROCESSING COMPLETED FOR L13

2 DUP REM L13 (0 DUPLICATES REMOVED)

=> d tot ibib abs

L14 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER:

2002:657919 HCAPLUS

DOCUMENT NUMBER:

137:195593

TITLE: Methods for the treatment of neuropathic pain by aryl

nitrone compounds

```
INVENTOR(S):
                        Waterbury, David; Wood, Paul L.; Khan, M. Amin;
                        Upasani, Ravindra B.
PATENT ASSIGNEE(S):
                        Centaur Pharmaceuticals, Inc., USA
SOURCE:
                        PCT Int. Appl., 82 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                    KIND DATE
                                         APPLICATION NO. DATE
     -----
                                         -----
     WO 2002065993 A2 20020829
WO 2002065993 A3 20021107
                                         WO 2002-US758 20020108
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
            PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
            UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU,
            TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     US 2002165274
                     A1 20021107
                                        US 2002-43659 20020108
PRIORITY APPLN. INFO.:
                                      US 2001-260469P P 20010108
OTHER SOURCE(S):
                       MARPAT 137:195593
    Methods are disclosed for the treatment of neuropathic pain by aryl
     nitrone compds. Method involves administration of an effective
     neuropathic pain-treating dose of a pharmaceutical compn. (Markush
     structures are given). Substituted aryl nitrone compds. are useful as
     therapeutics for neuropathic pain conditions in mammals.
L14 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                        2000:384142 HCAPLUS
DOCUMENT NUMBER:
                        133:30572
TITLE:
                        Preparation of 3,4,5-trisubstituted aryl nitrones for
                        the treatment of inflammation-related conditions
INVENTOR(S):
                        Waterbury, L. David; Wilcox, Allan L.; Carney, John
                        M.; Mavandadi, Farah; Danielzadeh, Albert
PATENT ASSIGNEE(S):
                        Centaur Pharmaceuticals, Inc., USA
SOURCE:
                        PCT Int. Appl., 73 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
    PATENT NO.
                   KIND DATE
                                        APPLICATION NO. DATE
     -----
                                         -----
                                       WO 1999-US28479 19991201
    WO 2000032567
                   A1 20000608 ·
        W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
            CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
            IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA,
            MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,
            SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM,
            AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
            DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
            CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                A 20010821 BR 1999-15886
A1 20010926 EP 1999-962967
    BR 9915886
                                                          19991201
    EP 1135367
                                                          19991201
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO
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US 6342523

B1 20020129

US 1999-452529

19991201

JP 2002531435 **T2** 20020924 JP 2000-585209 19991201 NO 2001002727 Α 20010726 NO 2001-2727 20010601 US 2003078297 US 2002-196800 A1 20030424 20020715 PRIORITY APPLN. INFO.: US 1998-110541P A2 19981202 WO 1999-US28479 W 19991201 US 2001-857264 A1 20010907

OTHER SOURCE(S):

MARPAT 133:30572

GI

The title compds. (I) [wherein R1 = C(W)R5, C(W)NR6R7, or CHR9XR8; R2 = alkyl or cycloalkylalkyl; R3 = H, (cyclo)alkyl, or aryl; R4, R5, and R8 = independently (un)substituted (cyclo)alkyl, (cyclo)alkenyl, or alkynyl; R6, R7, and R9 = independently H or (un)substituted (cyclo)alkyl, (cyclo)alkenyl, or alkynyl; W = O or S; X = O, S, S(O), or SO2] were prepd. by condensing trisubstituted benzaldehydes with hydroxylamines. For example, reaction of 4-acetoxy-3,5-di-tert-butylbenzaldehyde with tert-butylhydroxylamine gave II (74%). In in vitro assays, II did not inhibit cyclooxygenase-I (COX-1) and cyclooxygenase-2 (COX-2). Representative invention compds. were tested in a no. of assays and were effective for reducing the induction of prostaglandin E2 (PGE2) and/or effective in the carrageenan, adjuvant, and/or collagen assay. I are useful in the treatment of arthritis and other inflammation-related conditions and as anal. reagents for detecting free radicals.

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> fil req COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 18.59 410.77 DISCOUNT AMOUNTS (FOR OUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -1.30-1.30

FILE 'REGISTRY' ENTERED AT 12:16:21 ON 21 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2003 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 20 JUL 2003 HIGHEST RN 551897-78-0 DICTIONARY FILE UPDATES: 20 JUL 2003 HIGHEST RN 551897-78-0

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

Uploading aryl nitrone for neuropathic pain.str

L15 STRUCTURE UPLOADED

=> d L15 HAS NO ANSWERS L15 STR

G1 0, S

Structure attributes must be viewed using STN Express query preparation.

=> s 115

SAMPLE SEARCH INITIATED 12:16:54 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 0 TO ITERATE

100.0% PROCESSED 0 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 0 TO 0 PROJECTED ANSWERS: 0 TO 0

L16 0 SEA SSS SAM L15

=> s l15 full FULL SEARCH INITIATED 12:17:03 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 13 TO ITERATE

100.0% PROCESSED 13 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.01

L17 0 SEA SSS FUL L15

=> fil stng COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 148.55 559.32 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -1.30

FILE 'STNGUIDE' ENTERED AT 12:17:34 ON 21 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY, JAPAN SCIENCE AND TECHNOLOGY CORPORATION, AND FACHINFORMATIONSZENTRUM KARLSRUHE

FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Jul 11, 2003 (20030711/UP).

=> fil medl hcapl biosis uspatf

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST 0.30 559.62

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE
ENTRY
SESSION
CA SUBSCRIBER PRICE

0.00
-1.30

FILE 'MEDLINE' ENTERED AT 12:20:35 ON 21 JUL 2003

FILE 'HCAPLUS' ENTERED AT 12:20:35 ON 21 JUL 2003
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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FILE 'BIOSIS' ENTERED AT 12:20:35 ON 21 JUL 2003 COPYRIGHT (C) 2003 BIOLOGICAL ABSTRACTS INC.(R)

FILE 'USPATFULL' ENTERED AT 12:20:35 ON 21 JUL 2003 CA INDEXING COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

=> s inflammat? or antiinflammat?
L18 751121 INFLAMMAT? OR ANTIINFLAMMAT?

=> s neuropath?
L19 114667 NEUROPATH?

=> s 118 and 119 L20 10692 L18 AND L19

=> s l18 (S) l19 L21 5107 L18 (S) L19

=> s l18 (S) neuropathic pain L22 880 L18 (S) NEUROPATHIC PAIN

=> s inflammation (S) neuropathic pain L23 331 INFLAMMATION (S) NEUROPATHIC PAIN

=> dup rem 123
PROCESSING COMPLETED FOR L23
L24 293 DUP REM L23 (38 DUPLICATES REMOVED)

=> s 124 not py>2001 L25 113 L24 NOT PY>2001

=> focus PROCESSING COMPLETED FOR L25 L26 113 FOCUS L25 1-

=> d ibib abs 1-5

L26 ANSWER 1 OF 113 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: 2002:26707 BIOSIS DOCUMENT NUMBER: PREV200200026707

TITLE: Opioids in chronic pain.

AUTHOR(S): Przewlocki, Ryszard (1); Przewlocka, Barbara

CORPORATE SOURCE: (1) Department of Molecular Neuropharmacology, Institute of

Pharmacology, 12 Smetna Street, 31-343, Krakow:

nfprzewl@cyf-kr.edu.pl Poland

SOURCE: European Journal of Pharmacology, (19 October, 2001) Vol.

429, No. 1-3, pp. 79-91. print.

ISSN: 0014-2999.

DOCUMENT TYPE:

General Review

LANGUAGE: English

The advance in our understanding of the biogenesis of various endogenous opioid peptides, their anatomical distribution, and the characteristics of the multiple receptors with which they interact open a new avenue for understanding the role of opioid peptide systems in chronic pain. The main groups of opioid peptides: enkephalins, dynorphins and beta-endorphin derive from proenkephalin, prodynorphin and proopiomelanocortin, respectively. Recently, a novel group of peptides has been discovered in the brain and named endomorphins, endomorphin-1 and -2. They are unique in comparison with other opioid peptides by atypical structure and high selectivity towards the mu-opioid receptor. Another group, which joined the endogenous opioid peptide family in the last few years is the pronociceptin system comprising the peptides derived from this prohormone, acting at ORL1 receptors. Three members of the opioid receptor family were cloned in the early 1990s, beginning with the mouse delta-opioid receptor (DOR1) and followed by cloning of mu-opioid receptor (MOR1) and kappa-opioid receptor (KOR1). These three receptors belong to the family of seven transmembrane G-protein coupled receptors, and share extensive structural homologies. These opioid receptor and peptide systems are significantly implicated in antinociceptive processes. They were found to be represented in the regions involved in nociception and pain. The effects of opioids in animal models of inflammatory pain have been studied in great detail. Inflammation in the periphery influences the central sites and changes the opioid action. Inflammation increased spinal potency of various opioid receptor agonists. In general, the antinociceptive potency of opioids is greater against various noxious stimuli in animals with peripheral inflammation than in control animals. Inflammation-induced enhancement of opioid antinociceptive potency is characteristic predominantly for mu opioid receptors, since morphine elicits a greater increase in spinal potency of mu- than of delta- and kappa-opioid receptor agonists. Enhancement of the potency of mu-opioid receptor agonists during inflammation could arise from the changes occurring in opioid receptors, predominantly in affinity or number of the mu-opioid receptors. Inflammation has been shown to alter the expression of several genes in the spinal cord dorsal horn. Several studies have demonstrated profound alterations in the spinal PDYN system when there is peripheral inflammation or chronic arthritis. Endogenous dynorphin biosynthesis also increases under various conditions associated with neuropathic pain following damage to the spinal cord and injury of peripheral nerves. Interestingly, morphine lacks potent analgesic efficacy in neuropathic pain. A vast body of clinical evidence suggests that neuropathic pain is not opioid-resistant but only that reduced sensitivity to systemic opioids is observed in this condition, and an increase in their dose is necessary in order to obtain adequate analgesia. Reduction of morphine antinociceptive potency was postulated to be due to the fact that nerve injury reduced the activity of spinal opioid receptors or opioid signal transduction. Our recent study with endogenous ligands of the mu-opioid receptor, endomorphins, further complicates the issue, since endomorphins appear to be effective in neuropathic pain. Identification of the involved differences may be of importance to the understanding of the molecular mechanism of opioid action in neuropathic pain, as well as to the development of better and more effective drugs for the treatment of neuropathic pain in humans.

L26 ANSWER 2 OF 113 HCAPLUS COPYRIGHT 2003 ACS on STN ACCESSION NUMBER: 1997:712313 HCAPLUS DOCUMENT NUMBER: 127:357434

TITLE: Peripheral modulatory effects of catecholamines in

inflammatory and neuropathic pain

AUTHOR(S): Raja, Srinivasa N.

CORPORATE SOURCE: Dep. Anesthesiol. Critical Care Med., Div. Pain Med.,

Johns Hopkins Hosp., Baltimore, MD, 21287-5354, USA

50

SOURCE: Advances in Pharmacology (San Diego) (1998),

42 (Catecholamines), 567-571

CODEN: ADPHEL; ISSN: 1054-3589

PUBLISHER: Academic

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review, with 9 refs. discussing the role played by the sympathetic nervous system in pathol. states assocd. with pain and hyperalgesia, such

as inflammation and neuropathic pain.

Topics include sympathetic efferents in normal tissue, role of sympathetics in inflammed tissues, role of sympathetics in neuropathic pain, peripheral sympathetic-somatic coupling after partial nerve injury, and pharmacol. of animal models of sympathetically maintained pain.

L26 ANSWER 3 OF 113 HCAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 2002:587155 HCAPLUS

DOCUMENT NUMBER: 138:130997

TITLE: Fluoxetine, a selective serotonin reuptake inhibitor

modulates inflammatory and neuropathic pain in the rat

Pal Singh, Vijay; Jain, Naveen K.; Kulkarni, S. K. AUTHOR (S):

CORPORATE SOURCE: Pharmacology Division, University Institute of

Pharmaceutical Sciences, Panjab University,

Chandigarh, 160 014, India

SOURCE: Inflammopharmacology (2001), 9(3), 219-228

CODEN: IAOAES; ISSN: 0925-4692

PUBLISHER: VSP BV DOCUMENT TYPE: Journal LANGUAGE: English

The clin. usefulness of classical tricyclic antidepressants has been indicated in a variety of neuropathic pain. The role of selective serotonin reuptake inhibitors (SSRIs) is, however, controversial in pain control. The present study was aimed at evaluating the efficacy of an SSRI, fluoxetine, in neuropathic pain involving

peripheral (carrageenan-induced inflammation) and central sensitization (spinal nerve ligation) in rats. Fluoxetine was also assessed for antinociceptive and antiphlogistic effect against acetic acid-induced chemonociception in mice and carrageenan-induced

inflammation. Fluoxetine (100-400 .mu.g, intraplantar administration) failed to attenuate either hyperalgesia or cold allodynia in any of the tests employed. Fluoxetine dose dependently increased paw vol. in the

absence or presence of an inflammatory stimulus which was not reversed by indomethacin (10 mg/kg, p.o). Fluoxetine was ineffective in reducing hyperalgesia and allodynia assocd. with the rat models. However, fluoxetine dose dependently decreased acetic acid-induced writhings. The

results indicated that 5-HT plays a differential role in pain modulation and may not be playing a major role in the maintenance of hyperalgesia and allodynia in the rat models. REFERENCE COUNT: 24

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 4 OF 113 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: 2001:88979 BIOSIS DOCUMENT NUMBER: PREV200100088979

TITLE: Gabapentin attenuates the inflammation-induced increase in

the release of neuropeptides from rat spinal cord slices.

AUTHOR (S): Fehrenbacher, J. C. (1); Eckerle, C.; Vasko, M. R.

CORPORATE SOURCE: (1) Indiana University School of Medicine, Indianapolis, IN

USA

SOURCE: Society for Neuroscience Abstracts, (2000) Vol. 26, No.

1-2, pp. Abstract No.-453.8. print.

Meeting Info.: 30th Annual Meeting of the Society of Neuroscience New Orleans, LA, USA November 04-09, 2000

Society for Neuroscience

. ISSN: 0190-5295.

DOCUMENT TYPE: Conference LANGUAGE: English SUMMARY LANGUAGE: English

Gabapentin reduces nociceptive behaviors in animal models of inflammation and relieves neuropathic pain in

humans but has minimal effects on acute pain tests. To ascertain whether gabapentin's antinociceptive actions could be mediated by an effect on sensory neurons, we examined whether this anticonvulsant could attenuate the effects of inflammation on release of immunoreactive substance P (iSP) and calcitonin-gene related peptide (iCGRP) from sensory terminals in the spinal cord. Unilateral inflammation was induced in Sprague-Dawley rats by injecting 150 mul of a Freund's adjuvant (CFA) solution into one hindpaw. Five days later, spinal cords were removed, divided midsagitally, chopped and superfused with Kreb'sbicarbonate buffer in the presence and absence of gabapentin. Peptide release was evoked with 500 nM capsaicin and iSP and iCGRP measured by radioimmunoassay. When injection of CFA produced all increase in paw size of 3.2 mm. inflammation resulted in an 2-fold increase in the capsaicin-evoked release of iSP and iCGRP from spinal cord compared to tissue from the noninflamed side. Exposing the spinal cord tissue to 10 muM gabapentin completely abolished the inflammation -induced augmentation of the capsaicin-evoked peptide release. Evoked release of iSP and iCGRP was reduced from 2.2 +- 0.6 to 0.4 +- 0.1 % of total content/9 min and from 5.4 +- 1.1 to 1.9 +- 0.3 % of total content 9 min. respectively. In contrast, treatment with either 10 muM or 100 muM gabapentin did not significantly alter capsaicin-evoked peptide release from spinal cord tissue of non-inflamed rats. These results support the notion that gabapentin alters the sensitivity of sensory neurons that occurs secondary to inflammation without affecting neurons under control conditions. These findings are consistent with the observations that gabapentin is effective in reducing inflammatory or neuropathic pain and not acute pain.

L26 ANSWER 5 OF 113 MEDLINE on STN ACCESSION NUMBER: 97112135 MEDLINE

DOCUMENT NUMBER: PubMed ID: 8953862 97112135

TITLE: N-methyl-D-aspartate (NMDA) receptor and pain.

AUTHOR: Yamamoto T

CORPORATE SOURCE: Department of Anesthesiology, School of Medicine, Chiba

University.

SOURCE: MASUI. JAPANESE JOURNAL OF ANESTHESIOLOGY, (1996 Nov) 45

(11) 1312-8. Ref: 24

Journal code: 0413707. ISSN: 0021-4892.

PUB. COUNTRY: Japan

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: Japanese

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199701

ENTRY DATE: Entered STN: 19970219

> Last Updated on STN: 19970219 Entered Medline: 19970121

AB It has been shown that an excitatory amino acid, such as glutamate and aspartate, plays an important role in the spinal nociceptive transmission. NMDA receptor is one of the receptors of excitatory amino acids. Glutamate is present in the terminals of small diameter primary afferent fibers, as well as in dorsal horn interneurons. It has been reported that NMDA receptor is not located postsynaptic to primary afferent input; rather it mediates excitation evoked by glutamate-releasing interneurons. Activation of chemosensitive afferents with chemical irritants generates a state of central sensitization in the spinal cord, and this hyperexcitability is blocked by NMDA antagonist. These data suggested that activation of chemosensitive afferents induces release of glutamate which activates NMDA receptor in dorsal horn interneurons, and that this NMDA receptor activation induces spinal sensitization. It has been suggested that this spinal sensitization plays an important role in the maintenance of neuropathic pain and hyperalgesia during inflammation. In the clinical trial, epidural administration of NMDA antagonist attenuated the level of allodynia in patients with postherpetic neuralgia. I think that spinal sensitization induced by NMDA receptor activation is the key mechanism to maintain neuropathic pain and hyperalgesia during inflammation.

=> s 126 and lupus

L27 4 L26 AND LUPUS

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L27 ANSWER 1 OF 4 USPATFULL on STN

ACCESSION NUMBER: 2001:52054 USPATFULL

TITLE: Substituted imidazoles useful in the treatment of

inflammatory diseases

INVENTOR(S): Beers, Scott A., Flemington, NJ, United States

Malloy, Elizabeth A., Flemington, NJ, United States Wachter, Michael P., Bloomsbury, NJ, United States

Wu, Wei, Somerville, NJ, United States

PATENT ASSIGNEE(S): Ortho-McNeil Pharmaceutical, Inc., Raritan, NJ, United

States (U.S. corporation)

RELATED APPLN. INFO.: Division of Ser. No. US 1998-62304, filed on 17 Apr

1998, now patented, Pat. No. US 5965583

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Fan, Jane

NUMBER OF CLAIMS: 9
EXEMPLARY CLAIM: 1
LINE COUNT: 997

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB This invention relates to a series of substituted imidazoles of Formula I ##STR1##

pharmaceutical compositions containing them and intermediates used in their manufacture. The compounds of the invention inhibit the production of a number of inflammatory cytokines, and are useful in the treatment of diseases associated with overproduction of inflammatory cytokines.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L27 ANSWER 2 OF 4 USPATFULL on STN

ACCESSION NUMBER: 2000:61584 USPATFULL

TITLE: Application of botulinum toxin to the management of

neurogenic inflammatory disorders

INVENTOR(S): First, Eric R., 52-N-St., South Boston, MA, United

States 02127

NUMBER DATE -----

PRIORITY INFORMATION:

US 1996-20400P

19960906 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility

PRIMARY EXAMINER:

Granted

Lankford, Jr., Leon B.

ASSISTANT EXAMINER:

Tate, Christopher R.

LEGAL REPRESENTATIVE: NUMBER OF CLAIMS:

Engellenner, Thomas J. Nutter, McClennen & Fish LLP

EXEMPLARY CLAIM:

LINE COUNT:

654

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides a method for the use of at least one serotype or a combination of serotypes of Botulinum neurotoxin either alone or in combination with other peptides or fusion proteins, that when administered in a safe and effective amount, antagonize and therefore decrease or block inflammation induced by the neurogenic mechanisms underlying or associated with inflammatory disorders, in particular, arthritis.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L27 ANSWER 3 OF 4 USPATFULL on STN

ACCESSION NUMBER: TITLE:

2000:34566 USPATFULL

2-substituted imidazoles useful in the treatment of

INVENTOR (S):

inflammatory diseases Beers, Scott A., Flemington, NJ, United States

Malloy, Elizabeth A., Flemington, NJ, United States Wachter, Michael P., Bloomsbury, NJ, United States

Wu, Wei, Somerville, NJ, United States

PATENT ASSIGNEE(S):

Ortho-McNeil Pharmaceutical, Inc., Raritan, NJ, United

States (U.S. corporation)

NUMBER KIND DATE -----

PATENT INFORMATION:

US 6040320

20000321

APPLICATION INFO.:

US 1998-106698 Utility

19980629 (9)

DOCUMENT TYPE: FILE SEGMENT:

Granted

PRIMARY EXAMINER:

Fan, Jane

LEGAL REPRESENTATIVE: Harbour, John

NUMBER OF CLAIMS:

10

EXEMPLARY CLAIM:

1

LINE COUNT:

1075

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention relates to substituted imidazoles of Formula I ##STR1## pharmaceutical compositions containing them, methods of using them and

intermediates useful in their manufacture. The compounds of the invention modulate the production of a number of inflammatory cytokines, and are useful in the treatment of diseases associated with the

production of inflammatory cytokines.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L27 ANSWER 4 OF 4 USPATFULL on STN

ACCESSION NUMBER:

1999:124919 USPATFULL

TITLE:

Substituted imidazoles useful in the treatment of

inflammatory disease

INVENTOR(S):

Beers, Scott A., Flemington, NJ, United States Malloy, Elizabeth A., Flemington, NJ, United States

Wachter, Michael P., Bloomsbury, NJ, United States

Wu, Wei, Somerville, NJ, United States

PATENT ASSIGNEE(S):

Ortho-McNeil Pharmaceutical, Inc., Raritan, NJ, United

States (U.S. corporation)

DATE NUMBER KIND -----

PATENT INFORMATION: US 5965583 19991012

APPLICATION INFO.: US 1998-62304 19980417 (9)

> NUMBER DATE -----

PRIORITY INFORMATION: US 1997-44252P 19970427 (60)

DOCUMENT TYPE: Utility FILE SEGMENT: Granted PRIMARY EXAMINER: Fan, Jane FILE SEGMENT: LEGAL REPRESENTATIVE: Harbour, John

NUMBER OF CLAIMS: 17 EXEMPLARY CLAIM: LINE COUNT: 1130

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention relates to a series of substituted imidazoles of Formula I ##STR1## pharmaceutical compositions containing them and intermediates used in their manufacture. The compounds of the invention inhibit the production of a number of inflammatory cytokines, and are useful in the treatment of diseases associated with overproduction of inflammatory cytokines.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> fil stng

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 26.23 585.85 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -1.30 -2.60

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FILE CONTAINS CURRENT INFORMATION. LAST RELOADED: Jul 11, 2003 (20030711/UP).

=> FIL MEDL HCAPL BIOSIS USPATF

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 0.18 586.03

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -2.60

FILE 'MEDLINE' ENTERED AT 12:26:03 ON 21 JUL 2003

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FILE 'USPATFULL' ENTERED AT 12:26:03 ON 21 JUL 2003 CA INDEXING COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

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=> s acute brachial neuritis
L28
            10 ACUTE BRACHIAL NEURITIS
=> s neuritis
L29
         15329 NEURITIS
=> s 129 and 118
L30
          2827 L29 AND L18
=> s 129 (s) 118
L31
          1349 L29 (S) L18
=> s 131 and 119
L32
          517 L31 AND L19
=> s 131 (S) 119
L33
           157 L31 (S) L19
=> dup rem 133
PROCESSING COMPLETED FOR L33
L34
            145 DUP REM L33 (12 DUPLICATES REMOVED)
=> s 134 and neuropathic pain
L35
            23 L34 AND NEUROPATHIC PAIN
=> focus
PROCESSING COMPLETED FOR L35
L36
             23 FOCUS L35 1-
=> d ibib abs 1-5
L36 ANSWER 1 OF 23 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                         2002:977582 HCAPLUS
DOCUMENT NUMBER:
                         138:37450
TITLE:
                         Ras-MEK-ERK1/2 signaling pathway in the production of
                         inflammatory and neuropathic pain
                         and uses for analgesic screening
                         Levine, Jon David; Messing, Robert O.
INVENTOR (S):
PATENT ASSIGNEE(S):
                         The Regents of the University of California, USA
SOURCE:
                         PCT Int. Appl., 134 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
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                            -----
     WO 2002102232
                      A2
                                           WO 2002-US19107 20020614
                            20021227
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
             RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,
             UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     US 2003008807
                       A1
                            20030109
                                           US 2002-173332
                                                           20020614
PRIORITY APPLN. INFO.:
                                        US 2001-298491P P 20010614
    This invention pertains to the discovery of a novel pathway that mediates
```

hyperalgesia, neuropathic pain, and inflammatory pain.

This pathway is a third independent pathway that involves activation of extracellular signal-regulated kinases (ERKs) 1 and 2. The pathway

comprises a Ras-MEK-ERK1/2 cascade that acts independent of PKA or PKC.epsilon. as a novel signaling pathway for the prodn. of inflammatory (and neuropathic) pain. This pathway presents numerous targets for a new class of analgesic agents.

L36 ANSWER 2 OF 23 USPATFULL on STN

ACCESSION NUMBER: 2003:30977 USPATFULL

TITLE: Method for treating neuropathic pain

and pharmaceutical preparation therefor

INVENTOR(S): Lavand'Homme, Patricia, Brussel, BELGIUM

APPLICATION INFO.: US 2002-141532 A1 20020507 (10)

NUMBER DATE

PRIORITY INFORMATION: US 2001-289063P 20010507 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: KNOBBE MARTENS OLSON & BEAR LLP, 2040 MAIN STREET,

FOURTEENTH FLOOR, IRVINE, CA, 92614

NUMBER OF CLAIMS: 18 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 3 Drawing Page(s)

LINE COUNT: 827

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention relates to a method for a sustained treatment and/or prophylaxis of neuropathic pain in mammal

comprising administering by peripheral nerve injection a neuropathic pain relieving composition comprising an alpha-2-adrenergic agonist.

The invention further relates to the use of an alpha-2-adrenergic agonist for the preparation of an injectable medicament for the sustained treatment and/or prophylaxis of neuropathic pain in mammal by peripheral nerve block.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L36 ANSWER 3 OF 23 USPATFULL on STN

ACCESSION NUMBER: 2000:50711 USPATFULL

TITLE: Treatment of neuropathic pain

INVENTOR(S): Fairbanks, Carolyn A., NE. Rochester, MN, United States

Wilcox, George L., N. Golden Valley, MN, United States

Laughin, Tinna M., Anoka, MN, United States

PATENT ASSIGNEE(S): Solvay Pharmaceuticals GmbH, Hannover, Germany, Federal

Republic of (non-U.S. corporation)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted

PRIMARY EXAMINER: Jones, Dwayne C.
ASSISTANT EXAMINER: Delacroix-Muirheid, C.

LEGAL REPRESENTATIVE: Evenson, McKeown, Edwards & Lenahan, P.L.L.C.

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 4 Drawing Figure(s); 4 Drawing Page(s)

LINE COUNT: 445

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method of relieving neuropathic pain uses

moxonidine and its physiologically compatible acid-addition salts for the treatment and/or prophylaxis of **neuropathic pain**. A composition comprising an effective amount of moxonidine, or a pharmaceutically acceptable salt thereof, in a pharmaceutically acceptable carrier, is administered to a subject in need of such treatment. The composition may be administered intrathecally.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L36 ANSWER 4 OF 23 USPATFULL on STN

ACCESSION NUMBER: 2002:295225 USPATFULL

TITLE: Use of aryl nitrone compounds in methods for treating

neuropathic pain

INVENTOR(S): Waterbury, L. David, San Carlos, CA, UNITED STATES

Wood, Paul L., Morgan Hill, CA, UNITED STATES Khan, M. Amin, Morgan Hill, CA, UNITED STATES Upasani, Ravindra B., San Jose, CA, UNITED STATES

NUMBER DATE

PRIORITY INFORMATION: US 2001-260469P 20010108 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: William H. Benz, BURNS, DOANE, SWECKER & MATHIS,

L.L.P., P.O. Box 1404, Alexandria, VA, 22313-1404

NUMBER OF CLAIMS: 3 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 6 Drawing Page(s)

LINE COUNT: 1813

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB 3,4,5-trisubstituted aryl nitrone compounds having the formula:

##STR1##

where R.sup.1--R.sup.4 are as defined in the specification are useful as therapeutics for **neuropathic pain** conditions in mammals.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L36 ANSWER 5 OF 23 USPATFULL on STN

ACCESSION NUMBER: 2003:11103 USPATFULL

TITLE: Novel signaling pathway for the production of

inflammatory pain and neuropathy

INVENTOR(S): Levine, Jon David, San Francisco, CA, UNITED STATES

Messing, Robert O., Foster City, CA, UNITED STATES

PATENT ASSIGNEE(S): The Regents of the University of California (U.S.

corporation)

NUMBER DATE

PRIORITY INFORMATION: US 2001-298491P 20010614 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: QUINE INTELLECTUAL PROPERTY LAW GROUP, P.C., P O BOX

458, ALAMEDA, CA, 94501

NUMBER OF CLAIMS: 141 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 29 Drawing Page(s)

LINE COUNT: 4135

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention pertains to the discovery of a novel pathway that mediates hyperalgesia, neuropathic pain, and inflammatory pain. This pathway is a third independent pathway that involves activation of extracellular signal-regulated kinases (ERKs) 1 and 2. The pathway comprises a Ras-MEK-ERK1/2 cascade that acts independent of PKA or PKC.epsilon. as a novel signaling pathway for the production of inflammatory (and neuropathic) pain.

This pathway presents numerous targets for a new class of analgesic agents.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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- L35 ANSWER 1 OF 23 HCAPLUS COPYRIGHT 2003 ACS on STN
- TI Ras-MEK-ERK1/2 signaling pathway in the production of inflammatory and neuropathic pain and uses for analgesic screening
- L35 ANSWER 2 OF 23 HCAPLUS COPYRIGHT 2003 ACS on STN
- TI Agmatine reverses pain induced by inflammation, neuropathy, and spinal cord injury
- L35 ANSWER 3 OF 23 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN TI The Effect of Neuritis and Inflammatory Substances on Slowly Conducting Afferent Fibers.
- L35 ANSWER 4 OF 23 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN TI Expression of neurotrophic factors in the dorsal root ganglion in a rat model of lumbar disc herniation.
- L35 ANSWER 5 OF 23 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
 TI Infiltration of immune cells and TNFalpha into the rat sciatic nerve in an experimental neuritis that evokes neuropathic pain.
- L35 ANSWER 6 OF 23 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN Neuropathic pain from an experimental neuritis of the rat sciatic nerve.
- L35 ANSWER 7 OF 23 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN TI Does a neuroimmune interaction contribute to the genesis of painful peripheral neuropathies.
- L35 ANSWER 8 OF 23 USPATFULL on STN .
- TI 1-(Adamantyl) amidines and their use in the treatment of conditions generally associated with abnormalities in glutamatergic transmission
- L35 ANSWER 9 OF 23 USPATFULL on STN
- TI Methods of treating cytokine mediated diseases
- L35 ANSWER 10 OF 23 USPATFULL on STN
- TI 1,4-disubstituted benzo-fused cycloalkyl urea compounds
- L35 ANSWER 11 OF 23 USPATFULL on STN
- TI Human ion channels
- L35 ANSWER 12 OF 23 USPATFULL on STN
- TI Methods of treating cytokine mediated diseases
- L35 ANSWER 13 OF 23 USPATFULL on STN

- TI 2-adamantanemethanamine compounds for treating abnormalities in glutamatergic transmission
- L35 ANSWER 14 OF 23 USPATFULL on STN
- TI Use of certain drugs for treating nerve root injury
- L35 ANSWER 15 OF 23 USPATFULL on STN
- TI Method for treating neuropathic pain and pharmaceutical preparation therefor
- L35 ANSWER 16 OF 23 USPATFULL on STN
- TI Treating pain by targeting hyperpolarization-activated, cyclic nucleotide-gated channels
- L35 ANSWER 17 OF 23 USPATFULL on STN
- TI Treating pain by targeting hyperpolarization-activated, cyclic nucleotide-gated channels
- L35 ANSWER 18 OF 23 USPATFULL on STN
- TI Novel signaling pathway for the production of inflammatory pain and neuropathy
- L35 ANSWER 19 OF 23 USPATFULL on STN
- TI 1-(adamantyl) amidines and their use in the treatment of conditions generally associated with abnormalities in glutamatergic transmission
- L35 ANSWER 20 OF 23 USPATFULL on STN
- TI Use of aryl nitrone compounds in methods for treating neuropathic pain
- L35 ANSWER 21 OF 23 USPATFULL on STN
- TI Capsaicin receptor ligands
- L35 ANSWER 22 OF 23 USPATFULL on STN
- TI Adamantanecarboximidamide derivatives and their use as NMDA antagonists
- L35 ANSWER 23 OF 23 USPATFULL on STN
- TI Treatment of neuropathic pain

=> d ibib abs 23

L36 ANSWER 23 OF 23 USPATFULL on STN

ACCESSION NUMBER:

2001:112371 USPATFULL

TITLE:

Adamantanecarboximidamide derivatives and their use as

NMDA antagonists

INVENTOR(S):

Monck, Nathaniel Julius Thomas, Berkshire, United

Kingdom

Gillespie, Roger John, Berkshire, United Kingdom Bird, Andrew James, Berkshire, United Kingdom

PATENT ASSIGNEE(S):

Vernalis Research Limited, Wokingham, United Kingdom

date

20000913 PCT 102(e) date

(non-U.S. corporation)

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 6262106	B1	20010717	
	WO 9938841		19990805	
APPLICATION INFO.:	US 2000-600168		20000913	(9)
	WO 1999-GB321		19990101	
			20000913	PCT 371

NUMBER DATE

PRIORITY INFORMATION:

GB 1998-2225

19980202

DOCUMENT TYPE: Utility
FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Barts, Sam

PRIMARY EXAMINER: Barts, Samuel LEGAL REPRESENTATIVE: Foley & Lardner

NUMBER OF CLAIMS: 15 EXEMPLARY CLAIM: 1 LINE COUNT: 809

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB ##STR1##

The compound of the above formula wherein R1-R5 are independently selected from hydrogen, alkyl and aryl and prodrugs thereof; and pharmaceutically acceptable salts thereof; and use of the compounds in therapy, particularly for treatment of a condition generally associated with abnormalities in glutamtergic transmission.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> s neuralgia

L37 14916 NEURALGIA

=> s 127 and 118

L38 4 L27 AND L18

=> d

L38 ANSWER 1 OF 4 USPATFULL on STN

AN 2001:52054 USPATFULL

TI Substituted imidazoles useful in the treatment of inflammatory diseases

IN Beers, Scott A., Flemington, NJ, United States Malloy, Elizabeth A., Flemington, NJ, United States Wachter, Michael P., Bloomsbury, NJ, United States Wu, Wei, Somerville, NJ, United States

PA Ortho-McNeil Pharmaceutical, Inc., Raritan, NJ, United States (U.S. corporation)

PI US 6214830 B1 20010410 AI US 1999-295156 19990420 (9)

RLI Division of Ser. No. US 1998-62304, filed on 17 Apr 1998, now patented, Pat. No. US 5965583

DT Utility

FS Granted LN.CNT 997

INCL INCLM: 514/256.000
INCLS: 544/333.000

NCL NCLM: 514/256.000 NCLS: 544/333.000

IC [7]

ICM: C07D403-04

ICS: A61K031-506 514/256; 544/333

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> log h

COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
46.63 632.66

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL
ENTRY SESSION
CA SUBSCRIBER PRICE

-0.65
-3.25

SESSION WILL BE HELD FOR 60 MINUTES

ACCESSION NUMBER:

1977:499910 CAPLUS

DOCUMENT NUMBER:

7:-87:-99910-n

TITLE:

Pancreatic islet cell antibodies in diabetes mellitus correlated with the duration and type of diabetes,

coexistent autoimmune disease and

HLA type

AUTHOR(S):

Vaughan, H.; Irvine, W. J.; McCallum, C. J.; Gray, R. S.; Campbell, C. J.; Duncan, L. J. P.; Farquhar, J.;

Morris, P. J.

CORPORATE SOURCE:

SOURCE:

Dep. Endocrinol., R. Infirm., Edinburgh, Scot.

J. Endocrinol. (1977), 73(3), 40P(02)-41P

CODEN: JOENAK

DOCUMENT TYPE:

LANGUAGE:

Journal English

AB In both insulin-dependent and insulin-independent diabetics (IDD and IID, resp.) the prevalence of humoral pancreatic islet cell antibodies (ICA6) was dependent on the duration of diabetes and showed no correlation with the patient's age. The ICA6 persisted for longer in IDD with assocd. overt organ-specific autoimmune disease than in those without

such.

assocd. disease. The prevalence of HLA-B8 was increased in ICA6-pos. patients (61%) compared with patients in whom ICA6 was neg. within 3 months of diagnosis (35%) and in controls (28%). HLA-B7

was

less prevalent in ICA6-pos. patients compared with controls. Autoimmunity to pancreatic islet cell may, therefore, provide a better basis for the classification of idiopathic diabetes than the age of onset or type of treatment required.

L36 ANSWER 36 OF 47 CAPLUS COPYRIGHT 1999 ACS

ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

I.

AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE:

LANGUAGE:

GI

CAPLUS 85 579

Ring-chain isomerism of N-(1-carboxyalkyl)nitrones.

C-Aryl-N-(1-carboxylalkyl)nitrones

Kliegel, Wolfgang; Graumann, Juergen Inst. Pharm. Chem., Tech. Univ. Braunschweig,

Braunschweig, D-3300, Fed. Rep. Ger. Liebigs Ann. Chem. (1984), (9), 1545-62

CODEN: LACHDL; ISSN: 0170-2041

Journal German

RCH:N+(O-)CR1R2CO2H [I; R = (un) substituted Ph; R1 = H, Me, Ph; R2 = H, AB Me, Et; R1R2 = (CH2)4, (CH2)5] were prepd. by alkylation of (Z)-RCH:NOHwith R1R2CBrCO2H or by condensation of HONHCR1R2CO2H with RCHO. The ring-chain isomerism between nitrone I and oxazolidine II could not be proven spectroscopically. Acylating I with (Ph2B)20 gave boron chelates III of the open-chain nitrone form, while acylating with carboxylic acids or isocyanates gave oxazolidones IV [R3 = Me, 3,5-(O2N)2C6H3, 3-ClC6H4NH].

Alkylating I with PhCOCH2Br gave esters RCH:N+(O-)CR1R2CO2CH2COPh.

IT. 17556-16-0P 86737-36-2P 93562-94-8P

93562-95-9P 93563-03-2P 93563-06-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)

(prepn. and cyclization of, with oxybis(diphenylborane))

RN 17556-16-0 CAPLUS

CN Glycine, N-[(4-methoxyphenyl)methylene]-, N-oxide (9CI) (CA INDEX NAME)

$$CH = N - CH_2 - CO_2H$$
MeO

RN 86737-36-2 CAPLUS
CN L-Alanine, N-[(4-methoxyphenyl)methylene]-, N-oxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

RN 93562-94-8 CAPLUS

CN Benzeneacetic acid, .alpha.-[[(4-methoxyphenyl)methylene]oxidoamino]-(9CI) (CA INDEX NAME)

$$CH = N - CH - CO_2H$$
MeO

RN 93562-95-9 CAPLUS

CN Glycine, N-[(2-methoxyphenyl)methylene]-, N-oxide (9CI) (CA INDEX NAME)

$$CH = N - CH_2 - CO_2H$$
OMe

RN 93563-03-2 CAPLUS

CN L-Alanine, N-[(2-methoxyphenyl)methylene]-, N-oxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry unknown.

RN 93563-06-5 CAPLUS
CN Glycine, N-[(4-hydroxy-3-methoxyphenyl)methylene]-, N-oxide (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{O} \\ \parallel \\ \text{HO} \end{array}$$

$$\begin{array}{c} \text{CH} & \text{C} \\ \text{M} \\ \text{O} \\ \text{OH} \end{array}$$

CAPLUS COPYRIGHT 1999 ACS 1987:515572 CAPLUS L36 ANSWER 32 OF 47

ACCESSION NUMBER:

1.00 1-1-15-5-72 DOCUMENT NUMBER:

TITLE: Intramolecular 1,3-dipolar cycloaddition reactions.

I.

Thermochemical reactivities and regioselectivities of

4-substituted phenyl (N-4-pentenyl) nitrones

AUTHOR (S): Chen, Qinghua; Meng, Min

CORPORATE SOURCE: Dep. Chem., Beijing Norm. Univ., Beijing, Peop. Rep.

SOURCE: Huaxue Xuebao (1986), 44(9), 927-33

> Journal Chinese

CODEN: HHHPA4; ISSN: 0567-7351

DOCUMENT TYPE:

LANGUAGE:

OTHER SOURCE(S): CASREACT 107:115572

$$R = \begin{pmatrix} 0 \\ C = N \\ CH_2 \end{pmatrix}$$

- Refluxing nitrones I [R = NO2 (II), Br (III), H (IV), MeO (V)] in toluene for 24 h gave 54-80% cycloadducts VI and VII (VI/VII = 2). The thermochem. reactivity is in the order of II > III > IV > V.
- IT 109830-20-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and cycloaddn. reaction of)

RN. 109830-20-8 CAPLUS

4-Penten-1-amine, N-[(4-methoxyphenyl)methylene]-, N-oxide, (Z)- (9CI) (CA INDEX NAME)

$$CH = N - (CH_2)_3 - CH = CH_2$$

$$CH = N - CH_2 - CH_2 - CH = CH - Me$$
MeO

CAPLUS COPYRIGHT 1999 ACS L36 ANSWER 31 OF 47

(1988:549392 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

1.01911-11419131912

TITLE:

Synthesis of 2-alkyl-3-aryloxaziridines

AUTHOR(S): Kloc, Krystian; Kubicz, Elzbieta; Mlochowski, Jacek;

Syper, Ludwik

CORPORATE SOURCE:

Inst. Org. Phys. Chem., Tech. Univ. Wroclaw, Wroclaw, PL-50-370, Pol.

SOURCE:

Synthesis (1987), (12), 1084-7 CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE: LANGUAGE:

Journal English

OTHER SOURCE(S):

CASREACT 109:149392

H.
$$C = N_0$$
 CMe3 H. N_0 CMe3

Imination of RCHO (R = p-tolyl, p-anisyl, m-HOC6H4, p-XC6H4, X = Cl, Br, 3,4-(MeO)2C6H3, p-Et2NC6H4, naphthyl, 2-furyl, pyridyl, etc.) with Me3CNH2

in the presence of mol. sieves gave 66-99% .apprx.20 azomethines I, which were oxidized with m-ClC6H4CO3H and Na2CO3 in CHCl3 to give 21-66% .apprx.20 oxaziridines II.

IT' 72995-53-0P

> RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)

(prepn. and spectra of)

RN 72995-53-0 CAPLUS

2-Propanamine, N-[(2,4-dimethoxyphenyl)methylene]-2-methyl-, N-oxide CN

(9CI)

(CA INDEX NAME)

L36 ANSWER 20 OF 47 CAPLUS COPYRIGHT 1999 ACS ACCESSION NUMBER: 1992 1933197 CAPLUS

DOCUMENT NUMBER:

116:193591

TITLE:

Synthesis and characterization of phenyl-substituted

C-phenyl-N-tert-butylnitrones and some of their

radical adducts

AUTHOR(S):

Hinton, Randall D.; Janzen, Edward G.

CORPORATE SOURCE:

Natl. Biomed. Cent. Spin Trapp. Free Radicals,

Oklahoma Med. Res. Found., Oklahoma City, OK, 73104,

USA

SOURCE:

J. Org. Chem. (1992), 57(9), 2646-51

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE:

Journal English

LANGUAGE:

Synthesis of C-phenyl-N-tert-butylnitrone (PBN) and several of its

with substituents in the 2-, 3-, or 4-position on the Ph ring is described. While a one-pot redn./condensation method proved suitable for most compds., it was necessary to prep. some examples by direct condensation or through oxidn. of the appropriate imine. The 1H NMR data for the 3-X- and 4-X-PBN's can be correlated with the Hammett equation. For the 3-X series .DELTA..delta. for the .alpha.-proton correlates best with .sigma.+ and has a correlation coeff. of 0.90. For the 4-X series a dual substituent parameter equation using .sigma.R0 gives the best correlation with r = 0.99. The hyperfine splitting consts. of the hydroxyl radical and hydroperoxyl radical adducts of several substituted PBN's are also included and their correlation with the Hammett equation

is

discussed.

IT 115995-22-7P 130995-65-2P

RL: PRP (Properties); SPN (Synthetic preparation); PREP
(Preparation)

reparation,

(prepn. and NMR of) RN 115995-22-7 CAPLUS

CN 2-Propanamine, N-{(3-methoxyphenyl)methylene}-2-methyl-, N-oxide (9CI) (CA INDEX NAME)

RN 130995-65-2 CAPLUS

N 2-Propanamine, N-[(2-methoxyphenyl)methylene]-2-methyl-, N-oxide (9CI) (CA INDEX NAME)

L36 ANSWER 11 OF 47 CAPLUS COPYRIGHT 1999 ACS

ACCESSION NUMBER: 1995:258379 CAPLUS

DOCUMENT NUMBER:

TITLE:

Synthesis and separation of the E and Z isomers of

simple aldonitrones

AUTHOR(S):

Sivasubramanian, Shanmugaperumal; Mohan, Ponnusamy;

Thirumalaikumar, Muniappan; Muthusubramanian,

Shanmugan

CORPORATE SOURCE:

Dep. Org. Chem., Madurai Kamaraj Univ., Madurai, 625

021, India

SOURCE:

J. Chem. Soc., Perkin Trans. 1 (1994), (23), 3353-4

CODEN: JCPRB4; ISSN: 0300-922X

DOCUMENT TYPE:

Journal .

LANGUAGE:

English ·

OTHER SOURCE(S):

CASREACT 122:160227

The uncommon E isomer of simple aldonitrones has been obtained in significant amts. for the first time in the case of .alpha.-phenyl-N-(.beta.-phenylethyl)nitrones.

=> d. hitstr 11

L36 ANSWER 11 OF 47 CAPLUS COPYRIGHT 1999 ACS

161325-26-4P 161325-27-5P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. and sepn. of the E and Z isomers of simple aldonitrones and isomerization and)

RN 161325-26-4 CAPLUS

CN Benzeneethanamine, N-[(4-methoxyphenyl)methylene]-, N-oxide, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 161325-27-5 CAPLUS

Benzeneethanamine, N-[(4-methoxyphenyl)methylene]-, N-oxide, (Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

1995:242557 CAPLUS

DOCUMENT NUMBER:

-1-22:::31-33:6

TITLE:

Preparation of

(S)-.alpha.-phenyl-2-pyridineethanamine

(S)-malate and its use as an anticonvulsant and in

treatment of neurodegenerative

disorders

INVENTOR(S):

Murray, Robert John; Mathisen, Donald; Balestra,

Michel

PATENT ASSIGNEE(S):

Fisons PLC, UK; Fisons Corp.

SOURCE:

PCT Int. Appl., 31 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PAT	TENT NO.		KI	ND	DATE		•	. 1	APPLI	CATI	ои ис	o. 	DATE				
٠.	 .WO	9422831		 A	1	1994	1013			VO 19	94-G	B651		1994	0329	•		
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	NO	9203006		Α		1992	0730		1	10 19	92-3	006		1992	0730			
	FI	9203540		Α		1992	0806		.]	FI 19	92-3	540		1992				
	WO	19320052		Α	1 .	1993	1014			NO 19	93-G	B689		1993	0401	•	.1	
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		9463802		Α	1	1994	1024			AU 19	94-6	3802		1994	0329			
	ÄU	682348.		В	2	1997	1002											
	·EP	691957		А	1	1996	0117			EP 19	94-9	1123	2	1994	0329			
	EP	691957		В	1	1997	0813										. D.M.	
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=>

AB (S)-.alpha.-phenyl-2-pyridineethanamine (S)-malate (I), useful in the treatment of neurodegenerative disorders and as an anticonvulsant, is prepd. by the pptn. from soln. of a mixt. of .alpha.-phenyl-2-pyridineethanamine or its salts (e.g., the hydrochloride)

and (S)-malic acid. Thus, I was prepd. and demonstrated a ED50 of 2.9 mg/kg (p.o.) in the prevention of hind limb tonic extension in rats induced by maximal electroshock.

ACCESSION NUMBER: DOCUMENT NUMBER:

1997:413439 CAPLUS

<u> 127: 80093</u>;

TITLE:

Screening for subclinical autoimmune thyroid diseases with highly sensitive assays for autoantibodies to thyroglobulin and thyroid

peroxidase

and serum thyrotropin concentrations Konno, Norimichi

AUTHOR(S):

Konno Endocr. Clin., Sapporo, 060, Japan

CORPORATE SOURCE: SOURCE:

Nippon Naibunpi Gakkai Zasshi (1997), 73(3), 451-461

CODEN: NNGZAZ; ISSN: 0029-0661

Nippon Naibunpi Gakki

PUBLISHER: DOCUMENT TYPE:

Journal

LANGUAGE:

Japanese

The prevalence of subclin. autoimmune thyroid diseases (AITD) was investigated with detns. of highly sensitive anti-thyroid peroxidase antibody (TPOAb), anti-thyroglobulin antibody (TgAb) and TSH in serum samples obtained from 2647 ostensibly healthy subjects (1887 males and

760

females, aged 46.0 yr, Mean) residing in Sapporo who were recruited at

the

hospital for medical examns. The prevalence of thyroid autoantibodies (TAA) was 9.1%, among which 23.6% was TgAb pos. only (>1.83 U/mL), while 46.3% was TPOAb pos. only (>0.34 U/mL), and the remaining 32.2% were both TPOAb and TgAb pos. TAA was pos. in 6.5% for males and 15.8% for females with an age-related increase. The prevalence of subclin. hyperthyroidism (TSH <0.15 mU/L) was 0.79%, of which 42.1% was pos. for TAA. The prevalence of subclin. hypothyroidism (TSH >5.0 mU/L) was 1.1%, of which 20% were TAA pos. The overall prevalence of subclin. autoimmune thyroiditis as defined by TAA pos. eu- and hypothyroidism was 8.8%; 6.3% for males and 15.5% for females. There was a significant inverse correlation between TPOAb and TSH levels (r = -0.21) in subjects with TPOAb pos. only and also a significant correlation between TgAb and TSH levels in subjects with pos. TgAb with or without TPOAb (r = 0.27). This study indicates that the combined anal. of TPOAb, TgAb and TSH may

provide more accurate information on the prevalence of AITD in the population study.

CAPLUS 1999:39839

DOCUMENT NUMBER:

130:217477

TITLE:

Immunosuppressive therapy of lupus nephritis

AUTHOR(S):

Dooley, M. A.; Falk, R. J.

CORPORATE SOURCE:

Department of Medicine, The University of North Carolina at Chapel Hill School of Medicine, Chapel

Hill, NC, USA

SOURCE:

Lupus (1998), 7(9), 630-634 CODEN: LUPUES; ISSN: 0961-2033

Stockton Press

DOCUMENT TYPE:

Journal; General Review

LANGUAGE:

PUBLISHER:

English

A review with 50 refs. Aggressive immunosuppressive therapy should be considered for patients with proliferative lupus nephritis as the risk for progression to end stage renal disease is high.

Intermittent

i.v. cyclophosphamide therapy improves renal survival; longer duration of therapy is assocd. with fewer relapse of nephritis and decreased risk of diminished renal function. While azathioprine therapy does not differ statistically from steroids alone in prolonging renal survival, this therapy may be considered in patients with few risk factors for progression to renal insufficiency. Methylprednisolone as a single therapy does not prolong renal survival compared with regimens including cyclophosphamide. Plasmapheresis remains under study but has not shown addnl. benefit in treatment of severe lupus nephritis. The

potential roles for cyclosporin A and mycophenylate mofetil in the

therapy

of proliferative lupus nephritis remain to be defined. Supportive care including rigorous control of hypertension, consideration of angiotensin receptor inhibition or blockade to reduce proteinuria and prolong renal function, control of hyperlipidemia, prevention of osteoporosis, and prevention of pregnancy remain important clin. goals. Current research efforts focus on genetic and socioeconomic factors involved in racial differences in expression of lupus nephritis, hormonal manipulation to preserve gonadal function during cyclophosphamide therapy, and the potential impact on lupus activity of estrogen-contg. oral contraceptives or postmenopausal hormone replacement therapy.

CCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

lesions.

AUTHOR (S):

Antoine;

CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE:

LANGUAGE: .

CD8

1999:17<u>79</u>24 CAPLUS

130::280743 Frequent enrichment of CD8 T cells reactive against common herpes viruses in chronic inflammatory

Towards a reassessment of the physiopathological. significance of T cell clonal expansions found in autoimmune inflammatory processes

Scotet, Emmanuel; Peyrat, Marie-Alix; Saulquin, Xavier; Retiere, Christelle; Couedel, Christelle; Davodeau, Francois; Dulphy, Nicolas; Toubert,

Bignon, Jean-Denis; Lim, Anick; Vie, Henri; Hallet, Marie-Martine; Liblau, Roland; Weber, Michel; Berthelot, Jean-Marie; Houssaint, Elisabeth; Bonneville, Marc

Inst. Biologie, Nantes, F-44035, Fr. Eur. J. Immunol. (1999), 29(3), 973-985 CODEN: EJIMAF; ISSN: 0014-2980

Wiley-VCH Verlag GmbH

Journal . English

The authors recently evidenced a dramatic enrichment for T cells reactive against Epstein-Barr virus (EBV) within inflamed joints of 2 rheumatoid arthritis patients. To assess the generality of this phenomenon and its relevance to autoimmunity, the authors studied the responses of CD8 T cells from patients with either acute or chronic inflammatory diseases (rheumatoid arthritis, ankylosing spondylitis, psoriatic arthritis, Reiter's syndrome, arthrosis, uveitis, multiple sclerosis, encephalitis) against viral proteins derived from EBV and another common herpes virus, human cytomegalovirus (CMV). T cell responses against EBV and/or CMV epitopes were frequently obsd. within

T cells derived from chronic inflammatory lesions, irresp. of their location (knee, eye, brain) and autoimmune features. In most cases, CD8

cells derived from affected organs yielded stronger antiviral T cell responses than CD8 T cells derived from patients' PBL, even in chronic inflammatory diseases devoid of autoimmune features or induced by defined bacterial agents. These results suggest that the presence of virus-specific T cells within inflamed lesions of patients suffering from autoimmune diseases is a general phenomenon assocd. with chronic inflammation rather than the initiating cause of the autoimmune process. Since this phenomenon was sometimes assocd. with long-term T repertoire biases within inflamed lesions, the physiopathol. significance of T cell clonal expansions found in a recurrent fashion within chronically inflamed autoimmune lesions should be interpreted with caution.

1999:291647 CAPLUS

DOCUMENT NUMBER:

130::291586

TITLE:

Rapamycin for treatment of cardiac

inflammatory disease

INVENTOR(S):

Armstrong, Jay Joseph

PATENT ASSIGNEE(S):

American Home Products Corporation, USA

Pat. Specif. (Aust.), 13 pp. SOURCE:

CODEN: ALXXAP

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE .
AU 695664	В2	19980820	AU 1996-47983	19960308
AU 9647983	A1 .	19960919		

US 1995-401742 PRIORITY APPLN. INFO.: A method is provided for treating cardiac inflammatory disease in a mammal which comprises administering rapamycin to the

mammal orally, parenterally, intravascularly, intranasally,

intrabronchially, transdermally, or rectally.

L10 ANSWER 1 OF 2 CAPLUS COPYRIGHT 1999 ACS

ACCESSION NUMBER: 1999:46416 CAPLUS

DOCUMENT NUMBER:

New Indications for thalidomide

AUTHOR (S):

De Jong-van den Berg, L. T. W.; Rutgers, J.; Cornel,

M. C.; Takx-Koehlen, B. C. M. J.

CORPORATE SOURCE:

Groningen Inst. Drug Studies, Rijksuniv. Groningen,

Groningen, 9713 AW, Neth.

SOURCE:

TITLE:

Ziekenhuisfarmacie (1998), 14(4), 190-193

CODEN: ZIFAEM; ISSN: 0169-2720

PUBLISHER:

Nederlandse Vereniging van Ziekenhuisapothekers

DOCUMENT TYPE: Journal; General Review

LANGUAGE:

Dutch

AB A brief review with 38 refs. is given on new indications for thalidomide. Besides the beneficial effects of thalidomide in erythema nodosum leprosy the drug is efficacious in other major disorders such as aphthae and ulcers in aids, tuberculosis, autoimmune diseases, and tumors.

L10 ANSWER 2 OF 2 CAPLUS COPYRIGHT 1999 ACS

ACCESSION NUMBER:

1993:579060 CAPLUS

DOCUMENT NUMBER:

119:179060

TITLE:

In vitro tumor necrosis factor production by

mononuclear cells from lepromatous leprosy patients and from patients with erythema nodosum leprosum Santos, Dilvani O.; Suffys, Philip N.; Bonifacio,

AUTHOR(S):

CORPORATE SOURCE:

Karla; Marques, Maria A.; Sarno, Euzenir N. Dep. Cell. Mol. Biol., Fed. Fluminense Univ.,

Valonguinho, 20400, Brazil

SOURCE:

Clin. Immunol. Immunopathol. (1993), 67(3, Pt. 1),

199-203

CODEN: CLIIAT; ISSN: 0090-1229

DOCUMENT TYPE:

LANGUAGE:

Journal English

AB The prodn. of tumor necrosis factor (TNF) by Mycobacterium leprae-stimulated phagocyte cells, isolated from lepromatous leprosy patients (LL) and normal individuals, was evaluated, using the highly TNF-sensitive mouse fibrosarcoma cell line WEHI164cll3. Mononuclear cells, isolated from all individuals studied, showed a low level of spontaneous TNF prodn., except for patients undergoing erythema nodosum leprosum (ENL), in which the authors found significantly higher levels of TNF. Addn. of M. leprae to the phagocyte cell culture enhanced TNF prodn.

in all groups studied, except in the group with untreated leprosy patients. Strongest M. leprae-induced TNF release was found in mononuclear cell cultures derived from ENL patients. Patients in the postreactional state showed significantly higher TNF levels than healthy controls. These findings support the idea that TNF plays a key role in the complex symptomatol. of ENL.

TITLE:

AUTHOR(S):

CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE:

LANGUAGE:

Unavailable

1995:549812 CAPLUS

Editorial: adrenal and gonadal autoimmune

diseases

Smith, Bernard Rees; Furmaniak, Jadwiga

FIRS Lab., RSR Ltd., Cardiff, UK
J. Clin. Endocrinol. Metab. (1995), 80(5), 1502-5

CODEN: JCEMAZ; ISSN: 0021-972X

Journal -English

L14 ANSWER 2 OF 821 CAPLUS COPYRIGHT 1999 ACS

ACCESSION NUMBER:

99:397013 CAPLUS

TITLE:

Effects of systemically applied allopurinol and prednisolone on experimental autoimmune

uveitis

AUTHOR (S):

Augustin, A. J.; Loeffler, K. U.; Sekundo, W.; Grus,

F. H.; Lutz, J.

CORPORATE SOURCE:

Department of Ophthalmology, University of Bonn,

Bonn,

D-53105, Germany

SOURCE:

Graefe's Arch. Clin. Exp. Ophthalmol. (1999), 237(6),

508-512

CODEN: GACODL; ISSN: 0721-832X Springer-Verlag

PUBLISHER:

Journal

DOCUMENT TYPE: LANGUAGE:

English

AB To compare the effects of allopurinol to those of prednisolone on the oxidative tissue damage and inflammatory response in exptl. autoimmune uveitis (EAU). Expts. were performed using 27 male Lewis rats. EAU was induced by means of crude retina ext., Freund's adjuvant and pertussis toxin. One group of animals served as controls and two groups were treated systemically, one with allopurinol and one with prednisolone.

At the end of the expts. lipid peroxides (LPO), myeloperoxidase activity (MPO), and histol. changes were detd. in the retinal tissue. LPO were measured by two different methods [thiobarbituric acid reactive

(TBARS) and malondialdehyde-like substances]. Allopurinol led to a significant redn. in LPO and MPO levels. The steroid treatment also resulted in a significant redn. in MPO activity but LPO were significantly

reduced only when measured as TBARS. Histol. changes were significantly reduced by allopurinol only. Allopurinol is more effective than prednisolone in treating EAU. Its efficacy can be explained by the antioxidative/antiinflammatory and probably immunol. action. The antiinflammatory effects of prednisolone are not sufficient to reduce the tissue damage. Allopurinol promises to be a useful alternative to steroids in the treatment of uveitis.

	Type	Hits	Search Text	DBs
1	BRS	2	"20030078297"	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
2	BRS	2	6258852.pn.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
3	BRS	2	6083989.pn.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
4	BRS	131	flitter	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
5	BRS	72742	neuropathic or pain	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
6	BRS	24	flitter and (neuropathic or pain)	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
7	BRS	8976	neuropathic or neuroleptic	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
8	BRS	31	flitter and (neuropathic or neuroleptic)	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
9	BRS	9	(flitter and (neuropathic or neuroleptic)) and nitrone	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
10	BRS	2	5665732.pn.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
11	BRS	61	aryl near nitrone	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
12	BRS	2458	neuropathic	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
13	BRS	3	(aryl near nitrone) and neuropathic	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
14	BRS	2	6342523.pn.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
15	BRS	194	514/579.ccls.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
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17	BRS	435	514/643.ccls.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
18	BRS	123	514/715.ccls.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB

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	Туре	Hits	Search Text	DBs
19	BRS	230	514/717.ccls.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
20	BRS	116	514/720.ccls.	USPAT; EPO; JPO; DERWENT; IBM_TDB
21	BRS	1	569/50	USPAT; US-PGPUB; EPO JPO; DERWENT; IBM_TDB
22	BRS	0	569/50.ccls.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
23	BRS	1355	514/579.ccls. or 514/642.ccls. or 514/643.ccls. or 514/715.ccls. or 514/717.ccls. or 514/720.ccls.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
24	BRS	4900	(neuropathic adj pain) or neuralgia	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM_TDB
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26	BRS	2	5455272.pn.	USPAT; US-PGPUB; EPO; JPO; DERWENT; IBM TDB

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	Time Stamp	Comments	Error Definition	Errors
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